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(54) Title: THE MANUFACTURE AND EXPRESSION OF LARGE STRUCTURAL GENES

#### (57) Abstract

Rapid and highly efficient procedures for the total synthesis of linear, double stranded DNA sequences in excess of about 200 base pairs in length, which sequences may comprise entire structural genes. Novel sequences are prepared from two or more DNA subunits provided with terminal regions comprising restriction endonuclease cleavage sites facilitating insertion of subunits into a selected vector for purposes of amplification during the course of the total assembly process. The total, finally-assembled sequences include at least one, and preferably two or more, unique restriction endonuclease cleavage site(s) at intermediate positions along the sequence, allowing for easy excision and replacement of subunits and the corresponding by facile preparation of multiple structural analogs of polypeptides coded for by the sequences. Manufactured genes preferably include codons selected from

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among alternative codons specifying the same amino acid on the basis of preferential expression in a projected host microorganism (e.g., E. coli) to be transformed. Illustrated is the preparation and expression of manufactured genes capable of directing synthesis of human immune and leukocyte interferons and of other biologically active proteinaceous products, which products differ from naturally-occurring forms in terms of the identity and/or relative position of one or more amino acids, and in terms of one or more biological and pharmacological properties but which substantially retain other such properties.

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THE MANUFACTURE AND EXPRESSION OF LARGE STRUCTURAL GENES

This is a continuation-in-part of co-pending U.S. Patent Application Serial No. 375,494, filed 5 May 6, 1982.

The present invention relates generally to the manipulation of genetic materials and, more particularly, to the manufacture of specific DNA sequences useful in recombinant procedures to secure the production of proteins of interest.

Genetic materials may be broadly defined as those chemical substances which program for and guide the manufacture of constituents of cells and viruses and direct the responses of cells and viruses.

- 15 A long chain polymeric substance known as deoxyribonucleic acid (DNA) comprises the genetic material
  of all living cells and viruses except for certain
  viruses which are programmed by ribonucleic acids
  (RNA). The repeating units in DNA polymers are four
- different nucleotides, each of which consists of either a purine (adenine or guanine) or a pyrimidine (thymine or cytosine) bound to a deoxyribose sugar to which a phosphate group is attached. Attachment of nucleotides in linear polymeric form is by means of fusion of
- 25 the 5' phosphate of one nucleotide to the 3' hydroxyl group of another. Functional DNA occurs in the form of stable double stranded associations of single strands of nucleotides (known as deoxyoligonucleotides), which associations occur by means of hydrogen bonding between



purine and pyrimidine bases [i.e., "complementary" associations existing either between adenine (A) and thymine (T) or guanine (G) and cytosine (C)]. By convention, nucleotides are referred to by the names of their constituent purine or pyrimidine bases, and the complementary associations of nucleotides in double stranded DNA (i.e., A-T and G-C) are referred to as "base pairs". Ribonucleic acid is a polynucleotide comprising adenine, guanine, cytosine and uracil (U), rather than thymine, bound to ribose and a phosphate group.

Most briefly put, the programming function of DNA is generally effected through a process wherein specific DNA nucleotide sequences (genes) are "trans-15 cribed" into relatively unstable messenger RNA (mRNA) polymers. The mRNA, in turn, serves as a template for the formation of structural, regulatory and catalytic proteins from amino acids. This translation process involves the operations of small RNA strands (tRNA) which transport and align individual amino 20 acids along the mRNA strand to allow for formation of polypeptides in proper amino acid sequences. mRNA "message", derived from DNA and providing the basis for the tRNA supply and orientation of any given 25 one of the twenty amino acids for polypeptide "expression", is in the form of triplet "codons" -- sequential groupings of three nucleotide bases. In one sense, the formation of a protein is the ultimate form of "expression" of the programmed genetic message provided 30 by the nucleotide sequence of a gene.

Certain DNA sequences which usually "precede" a gene in a DNA polymer provide a site for initiation of the transcription into mRNA. These are referred to as "promoter" sequences. Other DNA sequences, also usually "upstream" of (i.e., preceding) a gene in a given DNA polymer, bind proteins that determine



the frequency (or rate) of transcription initiation. These other sequences are referred to as "regulator" sequences. Thus, sequences which precede a selected gene (or series of genes) in a functional DNA polymer 5 and which operate to determine whether the transcription (and eventual expression) of a gene will take place are collectively referred to as "promoter/regulator" or "control" DNA sequences. DNA sequences which "follow" a gene in a DNA polymer and provide a signal for termination of the transcription into mRNA are referred to as "terminator" sequences.

A focus of microbiological processing for nearly the last decade has been the attempt to manufacture industrially and pharmaceutically significant 15 substances using organisms which do not intially have genetically coded information concerning the desired product included in their DNA. Simply put, a gene that specifies the structure of a product is either isolated from a "donor" organism or chemically synthesized and then stably introduced into another organism which is preferably a self-replicating unicellular microorganism. Once this is done, the existing machinery for gene expression in the "transformed" host cells operates to construct the desired product.

The art is rich in patent and literature 25 publications relating to "recombinant DNA" methodologies for the isolation, synthesis, purification and amplification of genetic materials for use in the transformation of selected host organisms. U.S. Letters Patent No. 4,237,224 to Cohen, et al., for example, relates to transformation of procaryotic unicellular host organisms with "hybrid" viral or circular plasmid DNA which includes selected exogenous DNA sequences. dures of the Cohen, et al. patent first involve manufac-35 ture of a transformation vector by enzymatically cleaving viral or circular plasmid DNA to form linear DNA



strands. Selected foreign DNA strands are also prepared in linear form through use of similar enzymes. The linear viral or plasmid DNA is incubated with the foreign DNA in the presence of ligating enzymes capable of effecting a restoration process and "hybrid" vectors are formed which include the selected foreign DNA segment "spliced" into the viral or circular DNA plasmid.

Transformation of compatible unicellular host organisms with the hybrid vector results in the formation of multiple copies of the foreign DNA in the host cell population. In some instances, the desired result is simply the amplification of the foreign DNA and the "product" harvested is DNA. More frequently, the goal of transformation is the expression by the host cells of the foreign DNA in the form of large scale synthesis of isolatable quantities of commercially significant protein or polypeptide fragments coded for by the foreign DNA. See also, e.g., U.S. Letters Patent Nos. 4,269,731 (to Shine), 4,273,875

The success of procedures such as described in the Cohen, et al. patent is due in large part to the ready availability of "restriction endonuclease" enyzmes which facilitate the site-specific cleavage of both the unhybridized DNA vector and, e.g., eukaryotic DNA strands containing the foreign sequences of interest. Cleavage in a manner providing for the formation of single stranded complementary "ends" on the double stranded linear DNA strands greatly enhances the likeli-30 hood of functional incorporation of the foreign DNA into the vector upon "ligating" enzyme treatment. A large number of such restriction endonuclease enzymes are currently commercially available [See, e.g., "BRL Restriction Endonuclease Reference Chart" appearing 35 in the "'81/'82 Catalog" of Bethesda Research Laboratories, Inc., Gaithersburg, Maryland.] Verification



of hybrid formation is facilitated by chromatographic techniques which can, for example, distinguish the hybrid plasmids from non-hybrids on the basis of molecular weight. Other useful verification techniques involve radioactive DNA hybridization.

Another manipulative "tool" largely responsible for successes in transformation of procaryotic cells is the use of selectable "marker" gene sequences.

Briefly put, hybrid vectors are employed which contain,

in addition to the desired foreign DNA, one or more DNA sequences which code for expression of a phenotypic trait capable of distinguishing transformed from non-transformed host cells. Typical marker gene sequences are those which allow a transformed procaryotic cell

to survive and propagate in a culture medium containing metals, antibiotics, and like components which would kill or severely inhibit propagation of non-transformed host cells.

Successful expression of an exogenous gene 20 in a transformed host microorganism depends to a great extent on incorporation of the gene into a transformation vector with a suitable promoter/regulator region present to insure transcription of the gene into mRNA and other signals which insure translation of the mRNA 25 message into protein (e.g., ribosome binding sites). It is not often the case that the "original" promoter/regulator region of a gene will allow for high levels of expression in the new host. Consequently, the gene to be inserted must either be fitted with a new, 30 host-accommodated transcription and translation regulating DNA sequence prior to insertion or it must be inserted at a site where it will come under the control of existing transcription and translation signals in the vector DNA.

It is frequently the case that the insertion of an exogenous gene into, e.g., a circular DNA plasmid



vector, is performed at a site either immediately following an extant transcription and translation signal or within an existing plasmid-borne gene coding for a rather large protein which is the subject of 5 high degrees of expression in the host. In the latter case, the host's expression of the "fusion gene" so formed results in high levels of production of a "fusion protein" including the desired protein sequence (e.g., as an intermediate segment which can be isolated by 10 chemical cleavage of large protein). Such procedures not only insure desired regulation and high levels of expression of the exogenous gene product but also result in a degree of protection of the desired protein product from attack by proteases endogenous to the 15 host. Further, depending on the host organism, such procedures may allow for a kind of "piggyback" transportation of the desired protein from the host cells into the cell culture medium, eliminating the need to destroy host cells for the purpose of isolating the desired 20 product.

while the foregoing generalized descriptions of published recombinant DNA methodologies may make the processes appear to be rather straightforward, easily performed and readily verified, it is actually the case that the DNA sequence manipulations involved are quite painstakingly difficult to perform and almost invariably characterized by very low yields of desired products.

As an example, the initial "preparation" of a gene for insertion into a vector to be used in transformation of a host microorganism can be an enormously difficult process, especially where the gene to be expressed is endogenous to a higher organism such as man. One laborious procedure practiced in the art is the systematic cloning into recombinant plasmids of the total DNA genome of the "donor" cells,



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generating immense "libraries" of transformed cells carrying random DNA sequence fragments which must be individually tested for expression of a product of interest. According to another procedure, total mRNA is isolated from high expression donor cells (presumptively containing multiple copies of mRNA coded for the product of interest), first "copied" into single stranded cDNA with reverse transcriptase enzymes, then into double stranded form with polymerase, and cloned. The procedure again generates a library 10 of transformed cells somewhat smaller than a total genome library which may include the desired gene copies free of non-transcribed "introns" which can significantly interfere with expression by a host microorganism. The above-noted time-consuming gene 15 isolation procedures were in fact employed in published recombinant DNA procedures for obtaining microorganism expression of several proteins, including rat proinsulin [Ullrich, et al., Science, 196, pp. 1313-1318 (1977)], human fibroblast interferon [Goedell, et al., Nucleic 20 Acids Research, 8, pp. 4087-4094 (1980)], mouse β-endorphin [Shine, et al., <u>Nature</u>, <u>285</u>, pp. 456-461 (1980)] and human leukocyte interferon [Goedell, et al., Nature, 287, pp. 411-416 (1980); and Goedell, et al., Nature, 290, pp. 20-26 (1981)]. 25

whenever possible, the partial or total manufacture of genes of interest from nucleotide bases constitutes a much preferred procedure for preparation of genes to be used in recombinant DNA methods. A requirement for such manufacture is, of course, knowledge of the correct amino acid sequence of the desired polypeptide. With this information in hand, a generative DNA sequence code for the protein (i.e., a properly ordered series of base triplet codons) can be planned and a corresponding synthetic, double stranded DNA segment can be constructed. A combination of manufac-



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turing and cDNA synthetic methodologies is reported to have been employed in the generation of a gene for human growth hormone. Specifically, a manufactured linear double stranded DNA sequence of 72 nucleotide base pairs (comprising codons specifying the first 24 amino acids of the desired 191 amino acid polypeptide) was ligated to a cDNA-derived double strand coding for amino acids Nos. 25-191 and inserted in a modified pBR322 plasmid at a locus controlled by a lac promotor/regulator sequence [Goedell, et al., Nature, 281, pp. 544-548 (1981)].

completely synthetic procedures have been employed for the manufacture of genes coding for relatively "short" biologically functional polypeptides, such as human somatostatin (14 amino acids) and human insulin (2 polypeptide chains of 21 and 30 amino acids, respectively).

In the somatostatin gene preparative procedure [Itakura, et al., Science, 198, pp. 1056-1063 (1977)] a 52 base pair gene was constructed wherein 42 base 20 pairs represented the codons specifying the required 14 amino acids and an additional 10 base pairs were added to permit formation of "sticky-end" single stran-. ded terminal regions employed for ligating the structural gene into a microorganism transformation vector. Specifically, the gene was inserted close to the end of a β-galactosidase enzyme gene and the resultant fusion gene was expressed as a fusion protein from which somatostatin was isolated by cyanogen bromide cleavage. Manufacture of the human insulin gene, as noted above, involved preparation of genes coding for a 21 amino acid chain and for a 30 amino acid chain. Eighteen deoxyoligonucleotide fragments were combined to make the gene for the longer chain, and eleven fragments were joined into a gene for the shorter chain. Each gene was employed to form a fusion gene



with a β-galactosidase gene and the individually expressed polypeptide chains were enzymatically isolated and linked to form complete insulin molecules. [Goedell, et al., Proc. Nat. Acad. Sci. U.S.A., 76, pp. 106-110 (1979).]

In each of the above procedures, deoxyoligonucleotide segments were prepared, and then sequentially ligated according to the following general procedure. [See, e.g., Agarwal, et al., Nature, 227, pp. 1-7 (1970) and Khorana, Science, 203, pp. 614-675 (1979)]. 10 An initial "top" (i.e., 5'-3' polarity) deoxyoligonucleotide segment is enzymatically joined to a second "top" segment. Alignment of these two "top" strands is made possible using a "bottom" (i.e., 3' to 5' polarity) 15 strand having a base sequence complementary to half of the first top strand and half of the second top strand. After joining, the uncomplemented bases of the top strands "protrude" from the duplex portion formed. A second bottom strand is added which includes 20 the five or six base complement of a protruding top strand, plus an additional five or six bases which then protrude as a bottom single stranded portion. The two bottom strands are then joined. Such sequential additions are continued until a complete gene sequence is developed, with the total procedure being very time-consuming and highly inefficient.

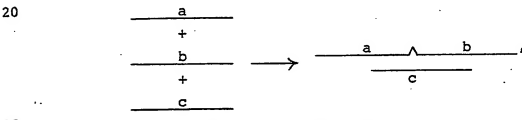
The time-consuming characteristics of such methods for total gene synthesis are exemplified by reports that three months' work by at least four investigators was needed to perform the assembly of the two "short", insulin genes previously referred to. Further, while only relatively small quantities of any manufactured gene are needed for success of vector insertion, the above synthetic procedures have such poor overall yields (on the order of 20% per ligation) that the eventual isolation of even minute quanti-



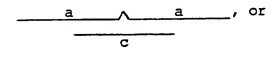
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ties of a selected short gene is by no means guaranteed with even the most scrupulous adherence to prescribed methods. The maximum length gene which can be synthesized is clearly limited by the efficiency with which the individual short segments can be joined. If n such ligation reactions are required and the yield of each such reaction is y, the quantity of correctly synthesized genetic material obtained will be proportional to y<sup>n</sup>. Since this relationship is expotential in nature, even a small increase in the yield per ligation reaction will result in a substantial increase in the length of the largest gene that may be synthesized.

Inefficiencies in the above-noted methodology are due in large part to the formation of undesired intermediate products. As an example, in an initial reaction forming annealed top strands associated with a bottom, "template" strand, the desired reaction may be,



25 but the actual products obtained may be



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or the like. Further, the longer the individual deoxyolidonucleotides are, the more likely it is that they will form thermodynamically stable self-associations such as "hairpins" or aggregations.



Proposals for increasing synthetic efficiency have not been forthcoming and it was recently reported that, "With the methods now available, however, it is not economically practical to synthesize genes for peptides longer than about 30 amino acid units, and many clinically important proteins are much longer".

[Aharonowitz, et al. Scientific American, 245, No. 3, pp. 140-152, at p. 151 (1981).]

An illustration of the "economic practicalities" involved in large gene synthesis is provided 10 by the recent publication of "successful" efforts in the total synthesis of a human leukocyte interferon gene. [Edge, et al., Nature, 292, pp. 756-782 (1981).] Briefly summarized, 67 different deoxyoligonucleotides 15 containing about 15 bases were synthesized and joined in the "50 percent overlap" procedure of the type noted above to form eleven short duplexes. in turn were assembled into four longer duplexes which were eventually joined to provide a 514 base pair gene coding for the 166 amino acid protein. The proce-20 dure, which the authors characterize as "rapid", is reliably estimated to have consumed nearly a year's effort by five workers and the efficiency of the assembly .strategy was clearly quite poor. It may be noted, for example, that while 40 pmole of each of the starting 67 deoxyoligonucleotides was prepared and employed to form the eleven intermediate-sized duplexes, by the time assembly of the four large duplexes was achieved, a yield of only about 0.01 pmole of the longer duplexes could be obtained for use in final assembly of the 30 whole gene.

Another aspect of the practice of recombinant DNA techniques for the expression, by microorganisms, of proteins of industrial and pharmaceutical interest is the phenomenon of "codon preference". While it was earlier noted that the existing machinery for



gene expression in genetically transformed host cells will "operate" to construct a given desired product, levels of expression attained in a microorganism can be subject to wide variation, depending in part on 5 specific alternative forms of the amino acid-specifying genetic code present in an inserted exogenous gene. A "triplet" codon of four possible nucleotide bases can exist in 64 variant forms. That these forms provide the message for only 20 different amino acids (as 10 well as transcription initiation and termination) means that some amino acids can be coded for by more Indeed, some amino acids have as than one codon. many as six "redundant", alternative codons while some others have a single, required codon. For reasons 15 not completely understood, alternative codons are not at all uniformly present in the endogenous DNA of differing types of cells and there appears to exist a variable natural hierarchy or "preference" for certain codons in certain types of cells.

As one example, the amino acid leucine is 20 specified by any of six DNA codons including CTA, CTC, CTG, CTT, TTA, and TTG (which correspond, respectively, to the mRNA codons, CUA, CUC, CUG, CUU, UUA and UUG). Exhaustive analysis of genome codon frequen-25 cies for microorganisms has revealed endogenous DNA of E. coli bacteria most commonly contains the CTG leucine-specifying codon, while the DNA of yeasts and slime molds most commonly includes a TTA leucinespecifying codon. In view of this hierarchy, it is generally held that the likelihood of obtaining high levels of expression of a leucine-rich polypeptide by an E. coli host will depend to some extent on the frequency of codon use. For example, a gene rich in TTA codons will in all probability be poorly expressed in E. coli, whereas a CTG rich gene will probably highly express the polypeptide. In a like manner,



when yeast cells are the projected transformation host cells for expression of a leucine-rich polypeptide, a preferred codon for use in an inserted DNA would be TTA. See, e.g., Grantham, et al. Nucleic Acids

5 Research, 8, pp. r49-r62 (1980); Grantham, et al., Nucleic Acids Research, 8, pp. 1893-1912 (1980); and, Grantham, et al., Nucleic Acids Research, 9, pp. r43-r74 (1981).

The implications of codon preference phenomena 10 on recombinant DNA techniques are manifest, and the phenomenon may serve to explain many prior failures to achieve high expression levels for exogenous genes in successfully transformed host organisms -- a less "preferred" codon may be repeatedly present in the 15 inserted gene and the host cell machinery for expression may not operate as efficiently. This phenomenon directs the conclusion that wholly manufactured genes which have been designed to include a projected host cell's preferred codons provide a preferred form of foreign 20 genetic material for practice of recombinant DNA techniques. In this context, the absence of procedures for rapid and efficient total gene manufacture which would permit codon selection is seen to constitute an even more serious roadblock to advances in the 25 art.

Of substantial interest to the background of the present invention is the state of the art with regard to the preparation and use of a class of biologically active substances, the interferons (IFNs).

Interferons are secreted proteins having fairly well-

Interferons are secreted proteins having fairly well-defined antiviral, antitumor and immunomodulatory characteristics. See, e.g., Gray, et al., Nature, 295, pp. 503-508 (1982) and Edge, et al., supra, and references cites therein.

On the basis of antigenicity and biological and chemical properties, human interferons have been



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grouped into three major classes: IFN-a (leukocyte), IFN- $\beta$  (fibroblast) and IFN- $\gamma$  (immune). Considerable information has accumulated on the structures and properties of the virus-induced acid-stable interferons 5 (IFN- $\alpha$  and  $\beta$ ). These have been purified to homogeneity and at least partial amino acid sequences have been determined. Analyses of cloned cDNA and gene sequences for IFN- $\beta_1$  and the IFN- $\alpha$  multigene family have permitted the deduction of the complete amino acid sequences of many of the interferons. In addition, efficient synthesis of IFN- $\beta_1$  and several IFN- $\alpha$ s in E. coli, and IFN- $\alpha_1$ , in yeast, have now made possible the purification of large quantities of these proteins in biologically active form.

Much less information is available concerning the structure and properties of IFN-y, an interferon . generally produced in cultures of lymphocytes exposed to various mitogenic stimuli. It is acid labile and does not cross-react with antisera prepared against IFN- $\alpha$  or IFN- $\beta$ . A broad range of biological activities have been attributed to IFN-y including potentiation of the antiviral activities of IFN- $\alpha$  and  $-\beta$ , from which it differs in terms of its virus and cell specificities and the antiviral mechanisms induced. In vitro studies performed with crude preparations suggest that the primary function of IFN-y may be as an immunoregulatory agent. The antiproliferative effect of IFN-y on transformed cells has been reported to be 10 to 100-fold greater than that of IFN- $\alpha$  or  $-\beta$ , suggesting a potential use in the treatment of neoplasia. Murine IFN-y preparations have been shown to have significant antitumor activity against mouse sarcomas.

It has recently been reported (Gray, et al., supra) that a recombinant plasmid containing a cDNA sequence coding for human IFN-y has been isolated and characterized. Expression of this sequence in



E. coli and cultured monkey cells is reported to give rise to a polypeptide having the properties of authentic human IFN-γ. In the publication, the cDNA sequence and the deduced 146 amino acid sequence of the "mature" polypeptide, exclusive of the putative leader sequence, is as follows:

10 Cys-Tyr-Cys-Gln-Asp-Pro-Tyr-Val-Lys-Glu-Ala-Glu-Asn-Leu-TGT TAC TGC CAG CAG CAA TAT GTA AAA GAA GCA GAA AAC CTT

20 Lys-Lys-Tyr-Phe-Asn-Ala-Gly-His-Ser-Asp-Val-Ala-Asp-Asn-AAG AAA TAT TTT AAT GCA GGT CAT TCA GAT GTA GCG GAT AAT

30 Gly-Thr-Leu-Phe-Leu-Gly-Ile-Leu-Lys-Asn-Trp-Lys-Glu-Glu-15 GGA ACT CTT TTC TTA GGC ATT TTG AAG AAT TGG AAA GAG GAG

50 Ser-Asp-Arg-Lys-Ile-Met-Gln-Ser-Gln-Ile-Val-Ser-Phe-Tyr-AGT GAC AGA AAA ATA ATG CAG AGC CAA ATT GTC TCC TTT TAC

60 /0
20 Phe-Lys-Leu-Phe-Lys-Asn-Phe-Lys-Asp-Asp-Gln-Ser-Ile-GlnTTC AAA CTT TTT AAA AAC TTT AAA GAT GAC CAG AGC ATC CAA

Lys-Ser-Val-Glu-Thr-Ile-Lys-Glu-Asp-Met-Asn-Val-Lys-Phe-AAG AGT GTG GAG ACC ATC AAG GAA GAC ATG AAT GTC AAG TTT

90
Phe-Asn-Ser-Asn-Lys-Lys-Arg-Asp-Asp-Phe-Glu-Lys-LeuTTC AAT AGC AAC AAA AAG AAA CGA GAT GAC TTC GAA AAG CTG

100 Thr-Asn-Tyr-Ser-Val-Thr-Asp-Leu-Asn-Val-Gln-Arg-Lys-Ala-30 ACT AAT TAT TCG GTA ACT GAC TTG AAT GTC CAA CGC AAA GCA

120
Ile-His-Glu-Leu-Ile-Gln-Val-Met-Ala-Glu-Leu-Ser-Pro-AlaATA CAT GAA CTC CTC ATC CAA ATG GCT GAA CTG TCG CAA GCA

130
Ala-Lys-Thr-Gly-Lys-Arg-Lys-Arg-Ser-Gln-Met-Leu-Phe-Gln-GCT AAA ACA GGG AAG CGA AAA AGG AGT CAG ATG CTG TTT CAA



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146 Gly-Arg-Arg-Ala-Ser-Gln GGT CGA AGA GCA TCC CAG.

In a previous publication of the sequence, arginine, rather than glutamine, was specified at position 140 in the sequence. (Unless otherwise indicated, therefore, reference to "human immune interferon" or, simply "IFN- $\gamma$ " shall comprehend both the [Arg<sup>140</sup>] and [Gln<sup>140</sup>] forms.)

activities of various interferon types makes the construction of synthetic polypeptide analogs of the interferons of paramount significance to the full development of the therapeutic potential of this class of compounds.

15 Despite the advantages in isolation of quantities of interferons which have been provided by recombinant DNA techniques to date, practitioners in this field have not been able to address the matter of preparation of synthetic polypeptide analogs of the interferons with any significant degree of success.

Put another way, the work of Gray, et al., supra, in the isolation of a gene coding for IFN-Y and the extensive labors of Edge, et al., supra, in providing a wholly manufactured IFN- $\alpha_1$  gene provide only genetic materials for expression of single, very precisely defined, polypeptide sequences. There exist no procedures (except, possibly, for site specific mutagenesis) which would permit microbial expression of large quantities of human IFN-y analogs which differed from the "authentic" polypeptide in terms of the identity or location of even a single amino acid. In a like manner, preparation of an IFN- $\alpha_1$  analog which differed by one amino acid from the polypeptide prepared by Edge, et al., supra, would appear to require an additional year of labor in constructing a whole new gene which varied in terms of a single triplet



codon. No means is readily available for the excision of a fragment of the subject gene and replacement with a fragment including the coding information for a variant polypeptide sequence. Further, modification of the reported cDNA-derived and manufactured DNA sequences to vary codon usage is not an available "option".

Indeed, the only report of the preparation of variant interferon polypeptide species by recombinant 10 DNA techniques has been in the context of preparation and expression of "hybrids" of human genes for IFN-α1 and IFN-α2 [Weck, et al., Nucleic Acids Research, 9, pp. 6153-6168 (1981) and Streuli, et al., Proc. Nat. Acad. Sci. U.S.A., 78, pp. 2848-2852 (1981)].

15 The hybrids obtained consisted of the four possible combinations of gene fragments developed upon finding that two of the eight human (cDNA-derived) genes fortuitously included only once within the sequence, base sequences corresponding to the restriction endonuclease cleavage sites for the bacterial endonucleases, PvuII and BglII.

There exists, therefore, a substantial need in the art for more efficient procedures for the total synthesis from nucleotide bases of manufactured DNA sequences coding for large polypeptides such as the interferons. There additionally exists a need for synthetic methods which will allow for the rapid construction of variant forms of synthetic sequences such as will permit the microbial expression of synthetic polypeptides which vary from naturally occurring forms in terms of the identity and/or position of one or more selected amino acids.



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### BRIEF SUMMARY

The present invention provides novel, rapid and highly efficient procedures for the total synthesis 5 of linear, double stranded DNA sequences in excess of about 200 nucleotide base pairs in length, which sequences may comprise entire structural genes capable of directing the synthesis of a wide variety of polypeptides of interest.

According to the invention, linear, double stranded DNA sequences of a length in excess of about 200 base pairs and coding for expression of a predetermined continuous sequence of amino acids within a selected host microorganism transformed by a selected. DNA vector including the sequence, are synthesized 15 by a method comprising:

(a) preparing two or more different, subunit, linear, double stranded DNA sequences of about 100 or more base pairs in length for assembly in a selected 20 assembly vector,

each different subunit DNA sequence prepared comprising a series of nucleotide base codons coding for a different continuous portion of said predetermined sequence of amino acids to be expressed,

one terminal region of a first of said subunits comprising a portion of a base sequence which provides a recognition site for cleavage by a first restriction endonuclease, which recognition site is entirely present either once or not at all in said 30 selected assembly vector upon insertion of the subunit therein,

one terminal region of a second of said subunits comprising a portion of a base sequence which provides a recognition site for cleavage by a second restriction endonuclease other than said first endonuclease, which recognition site is entirely present



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once or not at all in said selected assembly vector upon insertion of the subunit therein,

at least one-half of all remaining terminal regions of subunits comprising a portion of a recognition site (preferably a palindromic six base recognition site) for cleavage by a restriction endonuclease other than said first and second endonucleases, which recognition site is entirely present once and only once in said selected assembly vector after insertion of all subunits thereinto; and

(b) serially inserting each of said subunit

DNA sequences prepared in step (a) into the selected
assembly vector and effecting the biological amplification of the assembly vector subsequent to each insertion,
thereby to form a DNA vector including the desired
DNA sequence coding for the predetermined continuous
amino acid sequence and wherein the desired DNA sequence
assembled includes at least one unique, preferably
palindromic six base, recognition site for restriction
endonuclease cleavage at an intermediate position
therein.

The above general method preferably further includes the step of isolating the desired DNA sequence from the assembly vector preferably to provide one

25 of the class of novel manufactured DNA sequences having at least one unique palindromic six base recognition site for restriction endonuclease cleavage at an intermediate position therein. A sequence so isolated may then be inserted in a different, "expression"

30 vector and direct expression of the desired polypeptide by a microorganism which is the same as or different from that in which the assembly vector is amplified. In other preferred embodiments of the method: at least three different subunit DNA sequences are prepared in step (a) and serially inserted into said selected assembly vector in step (b) and the desired manufactured



DNA sequence obtained includes at least two unique palindromic six base recognition sites for restriction endonuclease cleavage at intermediate positions therein; the DNA sequence synthesized comprises an entire structural gene coding for a biologically active polypeptide; and, in the DNA sequence manufactured, the sequence of nucleotide bases includes one or more codons selected, from among alternative codons specifying the same amino acid, on the basis of preferential expression characteristics of the codon in said selected host microorganism.

Novel products of the invention include manufactured, linear, double stranded DNA sequences of a length in excess of about 200 base pairs and coding for the expression of a predetermined continuous sequence of amino acids by a selected host microorganism transformed with a selected DNA vector including the sequence, characterized by having at least one unique palindromic six base recognition site for restriction endonuclease cleavage at an intermediate position therein. Also included are polypeptide products of the expression by an organism of such manufactured sequences.

Illustratively provided by the present invention are novel manufactured genes coding for the synthesis of human immune interferon (IFN-γ) and novel
biologically functional analog polypeptides which
differ from human immune interferon in terms of the
identity and/or location of one or more amino acids.

Also provided are manufactured genes coding for synthesis of human leukocyte interferon of the F subtype
("LeIFN-F" or "IFN-αF") and analogs thereof, along
with consensus human leukocyte interferons.

DNA subunit sequences for use in practice

35 of the methods of the invention are preferably synthesized from nucleotide bases according to the methods



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disclosed in co-owned, concurrently-filed U.S. Patent Application Serial No. 375,493, by Yitzhak Stabinsky, entitled "Manufacture and Expression of Structural Genes" (Attorney's Docket No. 6250). Briefly summarized the general method comprises the steps of:

(1) preparing two or more different, linear, duplex DNA strands, each duplex strand including a double stranded region of 12 or more selected complementary base pairs and further including a top single stranded terminal sequence of from 3 to 7 selected bases at one end of the strand and/or a bottom single stranded terminal sequence of from 3 to 7 selected bases at the other end of the strand, each single stranded terminal sequence of each duplex DNA strand

single stranded terminal sequence of any other duplex
DNA strand prepared; and
(2) annealing each duplex DNA strand prepared
in step (1) to one or two different duplex strands
prepared in step (1) having a complementary single
stranded terminal sequence, thereby to form a single

comprising the entire base complement of at most one

continuous double stranded DNA sequence which has a duplex region of at least 27 selected base pairs including at least 3 base pairs formed by complementary association of single stranded terminal sequences of duplex DNA strands prepared in step (1) and which has from 0 to 2 single stranded top or bottom terminal regions of from 3 to 7 bases.

In the preferred general process for subunit manufacture, at least three different duplex DNA strands are prepared in step (1) and all strands so prepared are annealed concurrently in a single annealing reaction mixture to form a single continuous double stranded DNA sequence which has a duplex region of at least 42 selected base pairs including at least two non-adjacent sets of 3 or more base pairs formed by comple-



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mentary association of single stranded terminal sequences of duplex strands prepared in step (1).

The duplex DNA strand preparation step (1) of the preferred subunit manufacturing process preferably comprises the steps of:

- (a) constructing first and second linear deoxyoligonucleotide segments having 15 or more bases in a selected linear sequence, the linear sequence of bases of the second segment comprising the total complement of the sequence of bases of the first segment except that at least one end of the second segment shall either include an additional linear sequence of from 3 to 7 selected bases beyond those fully complementing the first segment, or shall lack a linear sequence of from 3 to 7 bases complementary to a terminal sequence of the first segment, provided, however, that the second segment shall not have an additional sequence of bases or be lacking a sequence of bases at both of its ends; and,
- (b) combining the first and second segments under conditions conducive to complementary association between segments to form a linear, duplex DNA strand.

The sequence of bases in the double stranded DNA subunit sequences formed preferably includes one or more triplet codons selected from among alternative codons specifying the same amino acid on the basis of preferential expression characteristics of the codon in a projected host microorganism, such as yeast cells or bacteria, especially E. coli bacteria.

Also provided by the present invention are improvements in methods and materials for enhancing levels of expression of selected exogenous genes in E. coli host cells. Briefly stated, expression vectors are constructed to include selected DNA sequences 35 upstream of polypeptide coding regions which selected sequences are duplicative of ribosome binding site



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sequences extant in genomic <u>E. coli</u> DNA associated with highly expressed endogenous polypeptides. A presently preferred selected sequence is duplicative of the ribosome binding site sequence associated with <u>E. coli</u> expression of outer membrane protein F ("OMP-F").

Other aspects and advantages of the present invention will be apparent upon consideration of the following detailed description thereof.

## DETAILED DESCRIPTION

As employed herein, the term "manufactured" as applied to a DNA sequence or gene shall designate a product either totally chemically synthesized by

15 assembly of nucleotide bases or derived from the biological replication of a product thus chemically synthesized. As such, the term is exclusive of products "synthesized" by cDNA methods or genomic cloning methodologies which involve starting materials which are of biological origin. Table I below sets out abbreviations employed herein to designate amino acids and includes IUPAC-recommended single letter designations.

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# TABLE I

	Amino Acid	Abbreviation	IUPAC Symbol
5	Alanine '	Ala	A
	Cysteine	Cys	С
	Aspartic acid	Asp	D
	Glutamic acid	Glu	E ·
	Phenylalanine	Phe	F
10	Glycine	Gly	G
	Histidine	His	H
	Isoleucine '	Ile	I
	Lysine	Lys	ĸ
	Leucine	Leu	L
15	Methionine	Met	M
	Asparagine	Asn	N
	Proline	Pro	P
	Glutamine	Gln	Q
	Arginine	Arg	R
20	Serine	Ser	S
	Threonine	Thr	T
	Valine	Val	V
	Tryptophan	Trp	W
	.Tyrosine	Tyr	<b>Y</b> .

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The following abbreviations shall be employed for nucleotide bases: A for adenine; G for guanine; T for thymine; U for uracil; and C for cytosine. For ease of understanding of the present invention, Table II and II below provide tabular correlations between the 64 alternate triplet nucleotide base codons of DNA and the 20 amino acids and transcription termination ("stop") functions specified thereby. In order to determine the corresponding correlations for RNA, U is substituted for T in the tables.



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TABLE II

	FIRST POSITION		SECOND 3	POSITION	Ī	THIRD POSITION
5		T	С	A	G	
		Phe	Ser	Tyr	Cys	T
		Phe	Ser	Tyr	Cys	С
	T	Leu	Ser	Stop	Stop	A
10		Leu	Ser	Stop	Trp	G
	•	Leu	Pro	His .	Arg	T
		Leu	Pro	His	Arg	С
	С	Leu	Pro	Gln	Arg	A
15		Leu	Pro	Gln	Arg	G
1.0		Ile	Thr	Asn	Ser	T
•		Ile	Thr	Asn	Ser	С
	<b>A</b>	Ile	Thr	Lys	Arg	A
		Met	Thr .	Lys ·	Arg	G
20		Val	Ala	Asp	Gly	T
		Val	Ala	Asp	Gly	С
	G	Val	Ala	Glu	Gly	<b>A</b> .
		Val	Ala	Glu	Gly	G
25	:					

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#### TABLE III

	Amin	o Acid	Specifying Codon(s)			
5	(A)	Alanine	GCT,	GCC, GCA, GCG		
	(C)	Cysteine	TGT,	TGC		
	(D)	Aspartic acid	GAT,	GAC		
	(E)	Glutamic acid	GAA,	GAG		
	(F)	Phenylalanine	TTT,	TTC		
10	(G)	Glycine	GGT,	GGC, GGA, GGG		
	(H)	Histidine	CAT,	CAC		
	(I)	Isoleucine	ATT,	ATC, ATA		
	(K)	Lysine	AAA,	AAG		
	(L)	Leucine	TTA,	TTG, CTT, CTC, CTA, CTG		
15	(M)	Methionine	ATG			
	(N)	Asparagine	AAT,	. AAC		
	(P)	Proline	CCT,	CCC, CCA, CCG		
	(Q)	Glutamine	CAA,	CAG		
	(R)	Arginine	CGT,	CGC, CGA, CGG, AGA, AGG		
20	(S)	Serine	TCT,	TCC, TCA, TCG, AGT, AGC		
	(T)	Threonine	ACT,	ACC, ACA, ACG		
	(V)	Valine	GTT,	GTC GTA, GTG		
	(W)	Tryptophan	TGG			
	(Y)	Tyrosine	TAC,	TAT		
25	STOP		TAA,	TAG, TGA		

A "palindromic" recognition site for restriction endonuclease cleavage of double stranded DNA is one which displays "left-to-right and right-to30 left" symmetry between top and bottom base complements, i.e., where "readings" of complementary base sequences of the recognition site from 5' to 3' ends are identical. Examples of palindromic six base recognition sites for restriction endonuclease cleavage include the
35 sites for cleavage by HindIII wherein top and bottom strands read from 5' to 3' as AAGCTT. A non-palindromic six base restriction site is exemplified by the site

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for cleavage by EcoP15, the top strand of which reportedly reads CAGCAG. The bottom strand base complement, when read 5' to 3' is CTGCTG. Essentially by definition, restriction sites comprising odd numbers of bases 5 (e.g., 5, 7) are non-palindromic. Certain endonucleases will cleave at variant forms of a site, which may be palindromic or not. For example, XhoII will recognize a site which reads (any purine)GATC(any pyrimidine) including the palindromic sequence AGATCT and the non-palindromic sequence GGATCT. Referring to the 10 previously-noted "BRL Restriction Endonuclease Reference Chart, " endonucleases recognizing six base palindromic . sites exclusively include BbrI, ChuI, Hin173, Hin91R, HinbIII, HinbIII, HindIII, HinfII, HsuI, BglII, StuI, RruI, ClaI, AvaIII, PvuII, SmaI, XmaI, EccI, SacII, SboI, SbrI, ShyI, SstII, TglI, AvrII, PvuI, RshI, RspI, XniI, XorII, XmaIII, BluI, MsiI, ScuI, SexI, SgoI, SlaI, SluI, SpaI, XhoI, XpaI, Bcel70, Bsul247, PstI, SalPI, XmaII, XorI, EcoRI, Rsh630I, SacI, SstI, SphI, BamHI, BamKI, BamNI, BamFI, BstI, KpnI, SalI, XamI, HpaI, XbaI, AtuCI, BclI, CpeI, SstIV, AosI, MstI, BalI, AsuII, and MlaI. Endonucleases which recognize only non-palindromic six base sequences exclusively include TthlllII, EcoPl5, AvaI, and AvrI. 25 Endonucleases recognizing both palindromic and nonpalindromic six base sequences include HaeI, HgiAI, Acyl, AosII, AsuIII, AccI, ChuII, HincII, HindII, MnnI, XhoII, HaeII, HinHI, NgoI, and EcoRI'.

Upon determination of the structure of a

desired polypeptide to be produced, practice of the
present invention involves: preparation of two or
more different specific, continuous double stranded
DNA subunit sequences of 100 or more base pairs in
length and having terminal portions of the proper

configuration; serial insertion of subunits into a
selected assembly vector with intermediate amplifica-



tion of the hybrid vectors in a selected host organism; use of the assembly vector (or an alternate, selected "expression" vector including the DNA sequence which has been manufactured from the subunits) to transform

5 a suitable, selected host; and, isolating polypeptide sequences expressed in the host organism. In its most efficient forms, practice of the invention involves using the same vector for assembly of the manufactured sequence and for large scale expression of the polypeptide. Similarly, the host microorganism employed for expression will ordinarily be the same as employed for amplifications performed during the subunit assembly process.

The manufactured DNA sequence may be provided

with a promoter/regulator region for autonomous control

of expression or may be incorporated into a vector

in a manner providing for control of expression by

a promoter/regulator sequence extant in the vector.

Manufactured DNA sequences of the invention may suitably

be incorporated into existing plasmid-borne genes

(e.g., β-galactosidase) to form fusion genes coding

for fusion polypeptide products including the desired

amino acid sequences coded for by the manufactured

DNA sequences.

In practice of the invention in its preferred forms, polypeptides produced may vary in size from about 65 or 70 amino acids up to about 200 or more amino acids. High levels of expression of the desired polypeptide by selected transformed host organisms is facilitated through the manufacture of DNA sequences which include one or more alternative codons which are preferentially expressed by the host.

Manufacture of double stranded subunit DNA sequences of 100 to 200 base pairs in length may proceed according to prior art assembly methods previously referred to, but is preferably accomplished by means



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of the rapid and efficient procedures disclosed in the aforementioned U.S. Application S.N. 375,493 by Stabinsky and used in certain of the following examples of actual practice of the present invention. 5 Briefly put, these procedures involve the assembly from deoxyoligonucleotides of two or more different, linear, duplex DNA strands each including a relatively long double stranded region along with a relatively short single stranded region on one or both opposing 10 ends of the double strand. The double stranded regions are designed to include codons needed to specify assembly of an initial, or terminal or intermediate portion of the total amino acid sequence of the desired polypeptide. Where possible, alternative codons preferentially 15 expressed by a projected host (e.g., E. coli) are employed. Depending on the relative position to be assumed in the finally assembled subunit DNA sequence, the single stranded region(s) of the duplex strands will include a sequence of bases which, when complemented 20 by bases of other duplex strands, also provide codons specifying amino acids within the desired polypeptide sequence.

Duplex strands formed according to this procedure are then enzymatically annealed to the one or two different duplex strands having complementary short, single stranded regions to form a desired continuous double stranded subunit DNA sequence which codes for the desired polypeptide fragment.

High efficiencies and rapidity in total

sequence assembly are augmented in such procedures
by performing a single annealing reaction involving
three or more duplex strands, the short, single stranded
regions of which constitute the base complement of
at most one other single stranded region of any other

duplex strand. Providing all duplex strands formed
with short single stranded regions which uniquely



complement only one of the single stranded regions of any other duplex is accomplished by alternative codon selection within the context of genetic code redundancy, and preferably also in the context of codon preferences of the projected host organism.

The following description of the manufacture of a hypothetical long DNA sequence coding for a hypothetical polypeptide will serve to graphically illustrate practice of the invention, especially in the context of formation of proper terminal sequences on subunit DNA sequences.

A biologically active polypeptide of interest is isolated and its amino acids are sequenced to reveal a constitution of 100 amino acid residues in a given continuous sequence. Formation of a manufactured gene for microbial expression of the polypeptide will thus require assembly of at least 300 base pairs for insertion into a selected viral or circular plasmid DNA vector to be used for transformation of a selected host organism.

A preliminary consideration in construction of the manufactured gene is the identity of the projected microbial host, because foreknowledge of the host allows for codon selection in the context of codon preferences of the host species. For purposes of this discussion, the selection of an <u>E</u>. <u>coli</u> bacterial host is posited.

A second consideration in construction of the manufactured gene is the identity of the projected

30 DNA vector employed in the assembly process. Selection of a suitable vector is based on existing knowledge of sites for cleavage of the vector by restriction endonuclease enzymes. More particularly, the assembly vector is selected on the basis of including DNA sequences providing endonuclease cleavage sites which will permit easy insertion of the subunits. In this regard,



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the assembly vector selected preferably has at least two restriction sites which occur only once (i.e., are "unique") in the vector prior to performance of any subunit insertion processes. For the purposes of this description, the selection of a hypothetical circular DNA plasmid pBR 3000 having a single EcoRI restriction site, i.e., -CTTAAG- and a single PvuII

restriction site, i.e., -CAGCTG-, is posited.

The amino acid sequence of the desired polypeptide is then analyzed in the context of determining availability of alternate codons for given amino acids (preferably in the context of codon preferences of the projected <u>E. coli</u> host). With this information in hand, two subunit DNA sequences are designed, preferably having a length on the order of about 150 base pairs -- each coding for approximately one-half of the total amino acid sequences of the desired polypeptide. For purposes of this description, the two subunits manufactured will be referred to as "A" and "B".

The methods of the present invention as applied to two such subunits, generally call for: insertion of one of the subunits into the assembly vector; amplification of the hybrid vector formed; and insertion of the second subunit to form a second hybrid including the assembled subunits in the proper sequence. Because the method involves joining the two subunits together in a manner permitting the joined ends to provide a continuous preselected sequence of bases coding for a continuous preselected sequence of amino acids, there exist certain requirements concerning the identity and sequence of the bases which make up the terminal regions of the manufactured subunits which will be joined to another subunit. Because the method calls for joining subunits to the assembly



vector, there exist other requirements concerning the identity and sequence of the bases which make up those terminal regions of the manufactured subunits which will be joined to the assembly vector. Because 5 the subunits are serially, rather than concurrently, inserted into the assembly vector (and because the methods are most beneficially practiced when the subunits can be selectively excised from assembled form to allow for alterations in selected base sequences therein), 10 still further requirements exist concerning the identity of the bases in terminal regions of subunits manufactured. For ease of understanding in the following discussion of terminal region characteristics, the opposing terminal regions of subunits A and B are respectively referred to as A-1 and A-2, and B-1 and B-2, viz: 15

B-2	B-1	A-2	A-1
В		A	

20 Assume that an assembly strategy is developed wherein subunit A is to be inserted into pBR3000 first, with terminal region A-1 to be ligated to the vector at the EcoRI restriction site. In the simplest case, the terminal region is simply provided with an EcoRI "sticky end", i.e., a single strand of four bases (-AATT- or -TTAA-) which will complement a single stranded sequence formed upon EcoRI digestion of pBR3000. This will allow ligation of terminal region A-1 to the vector upon treatment with ligase enzyme. Unless the single strand at the end of terminal region A-1 is preceded by an appropriate base pair (e.g., 3'-CTTAA-), the entire recognition site will not be reconstituted upon ligation to the vector. Whether or not the EcoRI recognition site is reconstituted upon ligation (i.e., whether or not there will be 0 or 1 EcoRI sites remain-35 ing after insertion of subunit A into the vector)



is at the option of the designer of the strategy. Alternatively, one may construct the terminal region A-1 of subunit A to include a complete set of base pairs providing a recognition site for some other endo-5 nuclease, hypothetically designated "XXX", and then add on portions of the EcoRI recognition site as above to provide an EcoRI "linker". To be of practical use in excising subunit A from an assembled sequence, the "XXX" site should not appear elsewhere in the 10 hybrid plasmid formed upon insertion. The requirement for construction of terminal region A-1 is, therefore, that it comprise a portion (i.e., all or part) of a base sequence which provides a recognition site for cleavage by a restriction endonuclease, which 15 recognition site is entirely present either once or not at all in the assembly vector upon insertion of the subunit.

Assume that terminal region B-2 of subunit
B is also to be joined to the assembly vector (e.g.,
20 at the single recognition site for PvuII cleavage
present on pBR3000). The requirements for construction
of terminal region B-2 are the same as for construction
of A-1, except that the second endonuclease enzyme
in reference to which the construction of B-2 is made
25 must be different from that with respect to which
the construction of A-1 is made. If recognition sites
are the same, one will not be able to separately excise
segments A and B from the fully assembled sequence.

The above assumptions require, then, that

30 terminal region A-2 is to be ligated to terminal region
B-1 in the final pBR3000 hybrid. Either the terminal
region A-2 or the terminal region B-1 is constructed
to comprise a portion of a (preferably palindromic
six base) recognition site for restriction endonuclease
35 cleavage by hypothetical third endonuclease "YYY"
which recognition site will be entirely present once



and only once in the expression vector upon insertion of all subunits thereinto, i.e., at an intermediate position in the assemblage of subunits. There exist a number of strategies for obtaining this result. 5 In one alternative strategy, the entire recognition site of "YYY" is contained in terminal region A-2 and the region additionally includes the one or more portions of other recognition sites for endonuclease cleavage needed to (1) complete the insertion of subunit 10 A into the assembly vector for amplification purposes, and (2) allow for subsequent joining of subunit A to subunit B. In this case, terminal region B-1 would have at its end only the bases necessary to link it to terminal region A-2. In another alternative, the entire "YYY" recognition site is included in terminal 15 region B-1 and B-1 further includes at its end a portion of a recognition site for endonuclease cleavage which is useful for joining subunit A to subunit B.

As another alternative, terminal region B-1 may contain at its end a portion of the "YYY" 20 recognition site. Terminal region A-2 would then contain the entire "YYY" recognition site plus, at its end, a suitable "linker" for joining A-2 to the · assembly vector prior to amplification of subunit A (e.g., a PvuII "sticky end"). After amplification of the hybrid containing subunit A, the hybrid would be cleaved with "YYY" (leaving a sticky-ended portion of the "YYY" recognition site exposed on the end of A-2) and subunit B could be inserted with its B-1 terminal region joined with the end of terminal region 30 A-2 to reconstitute the entire "YYY" recognition site. The requirement for construction of the terminal regions of all segments (other than A-1 and B-2) is that one or the other or both (i.e., "at least half") comprise a portion (i.e., include all or part) of a recognition site for third restriction endonuclease cleavage,



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which recognition site is entirely present once and only once (i.e., is "unique") in said assembly vector after insertion of all subunits thereinto. To generate a member of the class of novel DNA sequences of the invention, the recognition site of the third endonuclease should be a six base palindromic recognition site.

While a subunit "terminal region" as referred to above could be considered to extend from the subunit end fully halfway along the subunit to its center, as a practical matter the constructions noted would 10 ordinarily be performed in the final 10 or 20 bases. Similarly, while the unique "intermediate" recognition site in the two subunit assemblage may be up to three times closer to one end of the manufactured sequence 15 than it is to the other, it will ordinarily be located near the center of the sequence. If, in the above description, a synthetic plan was generated calling for preparation of three subunits to be joined, the manufactured gene would include two unique restriction enzyme cleavage sites in intermediate positions at least one of which will have a palindromic six base recognition site in the class of new DNA sequences of the invention.

The significant advantages of the abovedescribed process are manifest. Because the manufactured gene now includes one or more unique restriction endonuclease cleavage sites at intermediate positions along its length, modifications in the codon sequence of the two subunits joined at the cleavage site may be effected with great facility and without the need to re-synthesize the entire manufactured gene.

Following are illustrative examples of the actual practice of the invention in formation of manufactured genes capable of directing the synthesis 35 of: human immune interferon (IFN $\gamma$ ) and analogs thereof; human leukocyte interferon of the F subtype (INF-aF)



and analogs thereof; and, multiple consensus leukocyte interferons which, due to homology to IFN-αF can be named as IFN-αF analogs. It will be apparent from these examples that the gene manufacturing method-ology of the present invention provides an overall synthetic strategy for the truly rapid, efficient synthesis and expression of genes of a length in excess of 200 base pairs within a highly flexible framework allowing for variations in the structures of products to be expressed which has not heretofore been available to investigators practicing recombinant DNA techniques.

#### EXAMPLE 1

In the procedure for construction of synthetic 15 genes for expression of human IFNy a first selection made was the choice of E. coli as a microbial host for eventual expression of the desired polypeptides. Thereafter, codon selection procedures were carried out in the context of E. coli codon preferences enumer-20 ated in the Grantham publications, supra. A second selection made was the choice of pBR322 as an expression vector and, significantly, as the assembly vector to be employed in amplification of subunit sequences. 25 In regard to the latter factor, the plasmid was selected with the knowledge that it included single BamHI, HindIII, and SalI restriction sites. With these restriction sites and the known sequence of amino acids in human immune interferon in mind, a general plan for formation of three "major" subunit DNA sequences (IF-3, 30 IF-2 and IF-1) and one "minor" subunit DNA sequence (IF-4) was evolved. This plan is illustrated by Table IV below.



TABLE IV

IF-4 HpaI

		CTAG	
	CAG	GIC	
	i S	ACG	ч
10	TAC	ATG	
	T.C.T	ACA	
6	ATG	TAC,	
<del> </del>	ACA	CIT CAA GIG CAT TIT TCC CAT AGC TGT TAC, ACA ATG ACG GIC	
	i S	AGC	
	GTA	CAT	٠,
0	AGG	TCC	
	AAA	$T_iT$	_
	GTA	CAT	
H	CAC	GTG	
	GIT	CAA	
	ACG CAA	GTT	ک۔
a	ACG	TGC GTT CAA GTG CAT	2
	AGT	TCA	
	ACT	TGA	
L	Ø	H	

BamHI

-l l 2 3 4 Met Cys Tyr Cys Gln



ILE IV (cont'd.)

Ecori	Ħ	Gln		p Pr	o Ty	r Va	11 L)	(s G)	u Al	.a G]	u As	in Le	u Ly	's Ly	ß. T	Asp Pro Tyr Val Lys Glu Ala Glu Asn Leu Lys Lys Tyr Phe		Asn	
A A	A ATT CGA CAG GAT CCG GCT GTC CTA GGC	A CA	8 D	A GGC	G TAC		GTT À	AAG GAA TTC CTT		GCA GAA CGT CTT	GAA AAC CTT TTG		CTG AAA AAA GAC TTT TTT	A AA	T T	TAC TTATE AF	TTC AAC AAG TTG	သူ့ ညူ	
			Bam HI	HI									1	•					
20 Ala	20 Ala Gly His		Ser	Asp Val Ala Asp Asn	Val	Ala	Asp		30 Gly Thr	30 Thr	Leu	Leu Phe Leu	Leu	Gly	Ile	Gly Ile Leu Lys Asn Trp	Lys	Asn	Trp
		000						7.5						2 1					
GCA	GCA GGC C	CAC	TCC	GAC GTA GCT GAT CTG CAT CGA CTA	GTA	GCT	GAT	AAC	922 299	ACC TGG	GGC ACC CTG CCG TGG GAC	TTC CTG AAG GAC	CTG		ATC	CTA	AAA TTT	AAA AAC <sup>1</sup> 7 TTT TTG 1	TGG
			40 -						38 -					İ		T	36		
40 Lys	40 Lys Glu Glu		Ser	Asp Arg Lys Ile Met	Arg	Lys	Ile		Gln	50 Ser	50 Ser Gln Ile	Ile	53 Val	H	HindIII	ij			
			3.3	,						-									
AAA	AAA GAG GAA		202	GAC CGT AAG ATC ATG CAG	CGT	AAG	ATC	ATG	CAG	TCT	TCT CAA ATT	ATT	GTA	ت 1	A				
771	ראר		ו פ	יי זער	פרט	זיי	זטפ	100	ָרָ פַּדְרָנָ	202	32	דשעד	נט	3	٦				

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TABLE IV (cont'd.)

IF-2

ECORI

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Val Ser Phe Tyr Phe Lys Leu Phe Lys Asn Phe Lys Asp Asp Gln Ser Ile Gln CAG GAT CAA TCC CTA GTT AGG CTG TTC AAG AAC TTC AAA GAC GAC AAG TTC TTG AAG TTT CTG AAA CTG TTC 27-GTC CAT TCG AAG ATG AAG TTT, AATT

HindIII

Lys Ser Val Glu Thr Ile Lys Glu Asp Met Asn Val Lys Phe Phe Asn Ser Asn Lys Lys

AAC AAG AAG TTG TTC TTC GTA AAA TIC TIT AAC AGC CAT TIT AAG AAA TIG TCG GTA GAA ACT ATT AAG'GAG GAC ATG AAC TCG AAG AGC TTC,

Bglii Lys Arg Asp Asp Phe Glu Lys Leu Thr Asn Tyr Ser Val Thr 100

ຫ CTA CTG ACT AAC TAC TCT GTT ACA' GAC TGA TTG, ATG AGA CAA TGT GAT GAC TTC GAG AAA CTA, CTG AAG CTC TTT ပ္ပ် ဗိပ္ပံ AAA TTT

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TABLE IV (cont'd.)

IF-1

, E	FI &		H	00	
S C	G E		Met	ATG TAC	
G]	CTG ATC CAA GTT GAC TAG GTT CAA		130 Gly Lys Arg Lys Arg Ser Gln Met	AG	
Ile	ATC	4	er G	ည္က ည ၁၉	
G1u	CTG		S S	T. A.	
37 u	r cac gaa ca car car car car car car car car ca		i Ar	ဗ္ဗာ	
i S	AC O		Lγs	AAG	
e H	₽ <b>₽</b>		Arg	252	•
E II	r AT		S Y	AA	
Ala		ì	30 1y 1	2 2	
10 rg Lys	AAA TTT	ŧ.	ਜ਼ਿਲ੍ਹੇ ਮੁ	v g g	
110 Arg	TOS SCA		rh.	AC	
In 7	AG O		Lys	AAC	
. L	20		Ala	GCA	
ι Va	CAC		\1.a	ပ္တပ္တ	
110 Leu Asn Val Gln Arg Lys Ala Ile His Glu Glu Ile Gln Val	AA( TT(		Ser Pro Ala Ala Lys Thr	TCT CCT GCG GCA AAG ACT GGC AAA CGC AAG CGT AGC CAG ATG C AGA GGA CGC CGT TTC TGA CCG TTT GCG TTC GCA TCG GTC TAC G	
Leu	CTG GAC 14	ľ	. ผ	F 4 1	
105 Asp	TAT	LII		T T A	
. ~ .	GA C	Bglii	Let	CTG	
	စီပ		Glu Leu	GAA	
H	T C?		Ala	GCT GAA GGA CTT	
ECORI	A ATT CAG GCA GAT CTG AAC GTG CAG CGT AAA GCT ATT GTC CTC CTA GAC TTG CAC GTC GCA TTT CGA TAA		120 Met	ATG TAC	
	_				

	[-]
Salı	AGC
	TAG
Stop	TGA
146 Gln	CAG GTA 2
Ser	TCT
Ala	GCT
Arg	CGT
Arg	၁၅၁ ၁၅၁
140 Gln Gly	GGT CCA
140 Gln	CAG
140 Arg	CGT



The "minor" sequence (IF-4) is seen to include codons for the 4th through 1st (5'-TGT TAC TGC CAG) amino acids and an ATG codon for an initiating methionine [Met-1]. In this construction, it also includes additional bases to provide a portion of a control involved in an expression vector assembly from pBR 322 as described infra.

Alternative form of subunit IFN-1 for use in synthesis of a manufactured gene for  $[Arg^{140}]IFN\gamma$  included the codon 5'-CGT in place of 5'-CAG (for  $[Gln^{140}]$ ) at the codon site specifying the 140th amino acid.

The codon sequence plan for the top strand of the polypeptide-specifying portion total DNA sequence synthesized was as follows:

5'-TGT-TAC-TGC-CAG-GAT-CCG-TAC-GTT-AAG-GAA-GCA-GAA-AAC-CTG-AAA-AAA-TAC-TTC-AAC-GCA-GGC-CAC-TCC-GAC-GTA-GCT-GAT-AAC-GGC-ACC-CTG-TTC-CTG-GGT-ATC-CTA-AAA-AAC-CTG-GAT-AAC-GCT-GAC-CTG-AAG-ATC-ATG-CAG-TCT-CAA-ATT-GTA-AGC-GAA-TCC-GAC-CTG-AAG-ATC-ATG-CAG-TCT-AAA-ACC-GAT-GTA-AAC-TTC-AAA-CTG-TTC-AAG-AAC-TTC-AAA-GAC-GAT-CAA-TCC-ATC-CAG-AAG-AGC-GTA-GAA-ACT-ATT-AAG-GAG-GAC-ATG-AAC-GTA-AAA-TCC-TTT-AAC-AGC-AAC-AAG-AAG-AAA-CGC-GAT-GAC-TTC-GAG-AAA-CTG-ACT-AAC-TCT-GTT-ACA-GAT-CTG-AAC-GTG-CAG-CGT-AAA-GCT-ATT-CAC-GAA-CTG-ATC-CAA-GTT-ATG-GCT-GAA-CTG-TCT-CCT-GCG-GCA-AAG-ACT-GGC-AAA-CGC-AAG-CGT-AGC-CAG-ATG-CTG-TTT-CAG-[or CGT]-CGT-CGC-CGT-GCT-GCT-CCT-CAG-[or CGT]-CGT-CGC-CGT-GCT-TCT-CAG.

In the above sequence, the control sequence bases and the initial methionine-specifying codon is not illustrated, nor are termination sequences or sequences providing a terminal SalI restriction site. Vertical lines separate top strand portions attributable to each of the subunit sequences.



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The following example illustrates a preferred general procedure for preparation of deoxyoligonucleotides for use in the manufacture of DNA sequences of the invention.

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## EXAMPLE 2

Oligonucleotide fragments were synthesized using a four-step procedure and several intermediate washes. Polymer bound dimethoxytrityl protected nucleoside in a sintered glass funnel was first stripped of its 5'-protecting group (dimethoxytrity1) using 3% trichloroacetic acid in dichloromethane for 1-1/2 minutes. The polymer was then washed with methanol, 15 tetrahydrofuran and acetonitrile. The washed polymer was then rinsed with dry acetonitrile, placed under argon and then treated in the condensation step as follows. 0.5 ml of a solution of 10 mg tetrazole in acetonitile was added to the reaction vessel containing polymer. Then 0.5 ml of 30 mg protected nucleoside phosphoramidite in acetronitrile was added. This reaction was agitated and allowed to react for 2 minutes. The reactants were then removed by suction and the polymer rinsed with acetonitrile. This was followed by the oxidation step wherein 1 ml of a solution containing 0.1 molar I, in 2-6-lutidine/H,0/THF, 1:2:2, was reacted with the polymer bound oligonucleotide chain for 2 minutes. Following a THF rinse capping was done using a solution of dimethylaminopyridine (6.5 g 30 in 100 ml THF) and acetic anhydride in the proportion 4:1 for 2 minutes. This was followed by a methanol rinse and a THF rinse. Then the cycle began again with a trichloroacetic acid in CH2Cl2 treatment. The cycle was repeated until the desired oligonucleotide 35 sequence was obtained.



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The final oligonucleotide chain was treated with thiophenol dioxane, triethylamine 1:2:2, for 45 minutes at room temperature. Then, after rinsing with dioxane, methanol and diethylether, the oligonucleotide was cleaved from the polymer with concentrated ammonium hydroxide at room temperature. After decanting the solution from the polymer, the concentrated ammonium hydroxide solution was heated at 60°C for 16 hours in a sealed tube.

Each oligonucleotide solution was then extracted four times with 1-butanol. The solution was loaded into a 20% polyacrylamide 7 molar urea electrophoresis gel and, after running, the appropriate product DNA band was isolated.

Subunits were then assembled from deoxyoligonucleotides according to the general procedure for assembly of subunit IF-1.

Following the isolation of the desired 14 DNA segments, subunit IF-1 was constructed in the following manner:

- One nanomole of each of the DNA fragments excluding segment 13 and segment 2 which contain 5' cohesive ends, were subjected to 5'-phosphorylation;
- 2. The complementary strands of DNA, segments
  13 and 14, 11 and 12, 9 and 10, 7 and 8, 5 and 6,
  3 and 4 and 1 and 2 were combined together, warmed
  to 90° and slowly cooled to 25°;
- 3. The resulting annealed pairs of DNA were combined sequentially and warmed to 37° and slowly 30 cooled to 25°;
  - 4. The concentration of ATP and DTT in the final tube containing segments 1 thru 14 was adjusted to 150  $\mu$ M and 18 mM respectively. Twenty units of T-4 DNA ligase was added to this solution and the reaction was incubated at 4° for 18 hrs;



- 5. The resulting crude product was heated to 90° for 2 min. and subjected to gel filtration on Sephadex G50/40 using 10 mM triethyl ammonium bicarbonate as the eluent;
- 5 6. The desired product was purified, follow-ing 5' phosphorylation, using an 8% polyacrylamide-TBE gel.

Subunits IF-2, IF-3 and IF-4 were constructed in a similar manner.

of the complete human immune interferon gene from subunits IF-1, IF-2, IF-3, and IF-4; procedures for the growing, under appropriate nutrient conditions, of transformed <u>E. coli</u> cells, the isolation of human immune interferon from the cells, and the testing of biological activity of interferon so isolated.

# EXAMPLE 3

The major steps in the general procedure for assembly of the complete human IFNγ specifying genes from subunits IF-1, IF-2, and IF-3 are illustrated in Figure 1.

The 136 base pair subunit IF-1 was electro25 eluted from the gel, ethanol precipitated and resuspended
in water at a concentration of 0.05 pmol/µl. Plasmid
pBR322 (2.0 pmol) was digested with EcoRI and SalI,
treated with phosphatase, phenol extracted, ethanol
precipitated, and resuspended in water at a concentra30 tion of 0.1 pmol/µl. Ligation was carried out with
0.1 pmol of the plasmid and 0.2 pmol of subunit IF-1,
using T-4 DNA ligase to form hybrid plasmid pINT1.
E. coli were transformed and multiple copies of pINT1

35 The above procedure was repeated for purposes of inserting the 153 base pair subunit IF-2 to form

were isolated therefrom.



pINF2 except that the plasmid was digested with EcoRI and BglII. The 153 base pair IF-3 subunit was similarly inserted into pINT2 during manufacture of pINT3 except that EcoRI and Hind III were used to digest the plasmid.

An IF-4 subunit was employed in the construction of the final expression vector as follows: Plasmid PVvI was purchased from Stanford University, Palo Alto, California, and digested with PvuII. Using standard procedures, an EcoRI recognition site was 10 inserted in the plasmid at a PvuII site. Copies of this hybrid were then digested with EcoRI and HpaI to provide a 245 base pair sequence including a portion of the trp promoter/operator region. By standard procedures, IF-4 was added to the HpaI site in order to 15 incorporate the remaining 37 base pairs of the complete trp translational initiation signal and bases providing codons for the initial four amino acids of immune interferon (Cys-Tyr-Cys-Gln). The resulting assembly was then inserted into pINT3 which had been digested 20 with EcoRI and BamHI to yield a plasmid designated pINTy-trpI7.

E. coli cells containing pINTγ-trpI7 were grown on K media in the absence of tryptophan to an O.D.600 of 1. Indoleacrylic acid was added at a concentration of 20 μg per ml and the cells were cultured for an additional 2 hours at 37°C. Cells were harvested by centrifugation and the cell pellet was resuspended in fetal calf serum buffered with HEPES (pH 8.0). Cells were lysed by one passage through a French press at 10,000 psi. The cell lysate was cleared of debris by centrifugation and the supernatant was assayed for antiviral activity by the CPE assay ["The Interferon System" Stewart, ed., Springer-Verlag, N.Y., N.Y. (1981)]. The isolated product of expression was designated γ-1.



This example relates to a modification in the DNA sequence of plasmid pINTγ-trpI7 which facilitated the use of the vector in the trp promoter-controlled expression of structural genes coding for, e.g., analogs of IFN-γ and IFN-αF.

# EXAMPLE 4

Segment IF-4, as previously noted, had been 10 constructed to include bases coding for an initial methionine and the first four amino acids of IFN-y as well as 37 base pairs (commencing at its 5' end with a HpaI blunt end) which completed at the 3' end of a trp promoter/operator sequence, including a Shine 15 Delgarno ribosome binding sequence. It was clear that manipulations involving sequences coding IFN-Y analogs and for polypeptides other than IFN-y would be facilitated if a restriction site 3' to the entire trp promoter/operator region could be established. By way 20 of illustration, sequences corresponding to IF-4 for other genes could then be constructed without having to reconstruct the entire 37 base pairs needed to reconstitute the trp promoter/operator and would only require bases at the 5' end such as would facilitate. 25 insertion in the proper reading frame with the complete promoter/operator.

Consistent with this goal, sequence IF-4
was reconstructed to incorporate an XbaI restriction
site 3' to the base pairs completing the trp promoter/30 operator. The construction is shown in Table V below.



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## TABLE V

HpaI

AA CTA GTA CGC AAG TTC ACG TAA AAA GGG
TT GAT CAT GCG TTC AAG TGC ATT TTT CCC

XbaI -1 1 2 3 4 BamHI Met Cys Tyr Cys Gln

TAT CTA GAA ATG TGT TAC TGC CAG
ATA GAT CTT TAC ACA ATG ACG GTC CTAG

This variant form of segment IF-4 was inserted in pINTγ-trpI7 (digested with HpaI and BamHI) to generate plasmid pINTγ-TXb4 from which the IFN-γ-specifying gene could be deleted by digestion with XbaI and SalI and the entire trp promoter/operator would remain on the large fragment.

The following example relates to construction of structural analogs of IFN- $\gamma$  whose polypeptide structure differs from that of IFN- $\gamma$  in terms of the the identity of location of one or more amino acids.

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#### EXAMPLE 5

A first class of analogs of IFN-γ was formed which included a lysine residue at position 81 in place of asparagine. The single base sequence change needed to generate this analog was in subunit IF-2 of Table IV in segments 35 and 36. The asparagine-specifying codon, AAC, was replaced by the lysine-specifying codon, AAG. The isolated product of expression of such a modified DNA sequence [Lys 81] IFN-γ, was designated γ-10.

Another class of IFNy analogs consists of polypeptides wherein one or more potential glycosilation sites present in the amino acid sequence are deleted. More particularly, these consist of [Arg<sup>140</sup>]IFNy or [Gln<sup>140</sup>]IFNy wherein the polypeptide sequence fails to include one or more naturally occurring sequences, [(Asn or Gln)-(ANY)-(Ser or Thr)], which are known to provide sites for glycosilation of the polypeptide. One such sequence in IFNy spans positions 28 through 10 30, (Asn-Gly-Thr), another spans positions 101 through 103 (Asn-Tyr-Ser). Preparation of an analog according to the invention with a modification at positions 28-30 involved cleavage of plasmid containing all four IFN-y subunits with BamHI and HindIII to delete 15 subunit IF-3, followed by insertion of a variant of subunit IF-3 wherein the AAC codon for asparagine therein is replaced by the codon for glutamine, CAG. (Such replacement is effected by modification of deoxyoligonucleotide segment 37 to include CAG rather than 20 AAC and of segment 38 to include GTC rather than TTG. See Table IV.) The isolated product of expression of such a modified DNA sequence, [Gln<sup>28</sup>] IFN-y, was designated  $\gamma$ -12. Polypeptide analogs of this type would likely not be glycosilated if expressed in yeast 25 cells. Polypeptide analogs as so produced are not expected to differ appreciably from naturally-occurring IFNy in terms of reactivity with antibodies to the natural form, or in duration of antiproliferative or immunomodulatory pharmacological effects, but may display enhanced potency of pharmacological activity in one or more manner.

Other classes of IFNy analogs consists of polypeptides wherein the [Trp<sup>39</sup>] residue is replaced by [Phe<sup>39</sup>], and/or wherein one or more of the methionine residues at amino acid positions 48, 80, 120 and 137



are replaced by, e.g., leucine, and/or wherein cysteines at amino acid positions 1 and 3 are replaced by, e.g., serine or are completely eliminated. These last-mentioned analogs may be more easily isolated upon microbial expression because they lack the capacity for formation of intermolecular disulfide bridge formation.

Replacement of tryptophane with phenylalanine at position 39 required substitution for a TGG codon in subunit IF-3 with TTC (although TTT could also have been used), effected by modification of the deoxy-oligonucleotide segment 33 (TGG to TTC) and overlapping segment 36 (TGA to TAC) used to manufacture IF-3. [Phe<sup>39</sup>, Lys<sup>81</sup>]IFN-γ, the isolated product of expression of such a modified DNA sequence (which also included the above-noted replacement of asparagine by lysine at position 81) was designated γ-5.

In a like manner, replacement of one or more methionines at positions 48, 80, 120, and 137, respectively, involves alteration of subunit IF-3 20 (with reconstruction of deoxyoligonucleotides 31, 32 and 34), subunit IF-2 (with reconstruction of deoxyoligonucleotide segments 21 and 22); and subunit IF-1 (with reconstruction of deoxyoligonucleotide segments 7 and 10 and/or 3 and 4). An analog of IFN-y wherein 25 threonine replaced methionine at position 48 was obtained by modification of segment 31 in subunit IF-3 to delete the methionine-specifying codon ATG and replace it with an ACT codon. Alterations in segments 34 (TAC [Thr 48, to TGA) were also needed to effect this change. 30 Lys<sup>81</sup>]IFN-y, the isolated product of expression of such a modified DNA sequence (also including a lysinespecifying codon at position 81) was designated  $\gamma$ -6.

Replacement or deletions of cysteines at positions 1 and 3 involves only alteration of subunit 35 IF-4. As a first example, modifications in construction of subunit IF-4 to replace both of the cysteine-specifying codons at positions 1 and 3 (TGT and TGC, respec-



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tively) with the serine-specifying codon, TCT, required reconstruction of only 2 segments (see e and f of [Ser<sup>1</sup>, Ser<sup>3</sup>, Lys<sup>81</sup>]IFN-y, the isolated product of expression of the thus modified [Lys81] IFN-Y DNA sequence, was designated y-2. As another example, [Lys<sup>1</sup>, Lys<sup>2</sup>, Gln<sup>3</sup>, Lys<sup>81</sup>] IFN- $\gamma$ , designated  $\gamma$ -3, was obtained as an expression product of a modified construction of subunit IF-4 wherein codons AAA, AAA, and CAA respectively replaced TTG, TAC and TGC. Finally, [des-Cys<sup>1</sup>, des-Tyr<sup>2</sup>, des-Cys<sup>3</sup>, Lys<sup>81</sup>]IFN-y, designated  $\gamma$ -4, was obtained by means of modification of subunit IF-4 sections to 3'-ATC CAG-3' in the amino acid specifying region. It should be noted that the above modifications in the initial amino acid coding regions of 15 the gene were greatly facilitated by the construction of pINTy-TXb4 in Example 4 which meant that only short sequences with XbaI and BamHI sticky ends needed to be constructed to complete the amino terminal protein coding sequence and link the gene to the complete trp promoter. 20

Among other classes of IFN-y analog polypeptide provided by the present invention are those including polypeptides which differ from IFN-y in terms of amino acids traditionally held to be involved in secondary and tertiary configuration of polypeptides. As an example, provision of a cysteine residue at an intermediate position in the IFN-y polypeptide may generate a species of polypeptide structurally facilitative of formation of intramolecular disulfide bridges between amino terminal and intermediate cysteine residues such as found in IFN-a. Further, insertion or deletion of prolines in polypeptides according to the invention may alter linear and bending configurations with corresponding effects on biological activity. [Lys<sup>81</sup>, Cys<sup>95</sup>]IFN- $\gamma$ , designted  $\gamma$ -9, was isolated upon expression of a DNA sequence fashioned with 3'-AGC-5'



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replacing 3'-AAG-5' in sections 17 and 18 of subunit IF-2. A DNA sequence specifying [Cys<sup>95</sup>]IFN-γ (to be designated γ-11) is being constructed by the same general procedure. Likewise, a gene coding for [Cys<sup>95</sup>, Pro<sup>104</sup>]IFN-γ is under construction with the threonine-specifying codon ACA (section 15 of IF-2) being replaced by the proline-specifying codon CCA.

[Glu<sup>5</sup>]IFN-γ, to be designated γ-13, will result from modification of section 43 in subunit IF-3 to include the glutamate codon, GAA, rather than the aspartic acid specifying codon, GAT. Because such a change would no longer permit the presence of a BamHI recognition site at that locus, subunit IF-3 will likely need to be constructed as a composite subunit with the amino acid specifying portions of subunit IF-4, leaving no restriction site between xbaI and HindIII in the assembled gene. This analog of IFN-γ is expected to be less acid labile than the naturally-occurring form.

The above analogs having the above-noted tryptophane and/or methionine and/or cysteine replacements are not expected to differ from naturally-occurring IFN $\gamma$  in terms of reactivity with antibodies to the natural form or in potency of antiproliferative or immunomodulatory effect but are expected to have enhanced duration of pharmacological effects.

Still another class of analogs consists of polypeptides of a "hybrid" or "fused" type which include one or more additional amino acids at the end of the prescribed sequence. These would be expressed by DNA sequences formed by the addition, to the entire sequence coding for IFNy, of another manufactured DNA sequence, e.g., one of the subunits coding for a sequence of polypeptides peculiar to LeIFN-Con, described infra. The polypeptide expressed is expected to retain at least some of the antibody reactivity of naturally-occurring IFNy and to display some degree



of the antibody reactivity of LeIFN. Its pharmacological activities are expected to be superior to naturally-occurring IFN- $\gamma$  both in terms of potency and duration of action.

5 Table VI, below, sets forth the results of studies of antiviral activity of IFN-γ prepared according to the invention along with that of certain of the analogs tested. Relative antiviral activity was assayed in human HeLa cells infected with encephalo10 myocarditis virus (EMCV) per unit binding to a mono-clonal antibody to IFN-γ as determined in an immunoabsorbant assay.

# TABLE VI

15		
]	Interferon	Relative Antiviral <u>Activity</u>
	γ-1	1.00
	γ-4	0.60
	γ <b>-</b> 5	0.10
20 .	γ-6	0.06
	γ-10	0.51

The following example relates to modifications in the polypeptide coding region of the DNA sequences of the previous examples which serve to enhance the expression of desired products.

#### EXAMPLE 6

Preliminary analyses performed on the polypeptide products of microbial expression of manufactured DNA sequences coding for IFN-γ and analogs of IFN-γ revealed that two major proteins were produced in approximately equal quantities — a 17K form corresponding to the complete 146 amino acid sequence and a 12K form corresponding to an interferon fragment missing about 50 amino acids of the amino terminal. Review

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of codon usage in the manufactured gene revealed the likelihood that the abbreviated species was formed as a result of microbial translation initiation at the Met<sup>48</sup> residue brought about by the similarity 5 of base sequences 3' thereto to a Shine-Delgarno ribosome binding sequence. It thus appeared that while about half of the transcribed mRNA's bound to ribosomes only at a locus prior to the initial methionine, the other half were bound at a locus prior to the Met 48 codon. 10 In order to diminish the likelihood of ribosome binding internally within the polypeptide coding region, sections 33 and 34 of subunit IF-3 were reconstructed. More specifically, the GAG codon employed to specify a glutamate residue at position 41 was replaced by the 15 alternate, GAA, codon and the CGT codon employed to specify arginine at position 45 was replaced by the alternate, CGC, codon. These changes, effected during construction of the gene specifying the  $\gamma$ -6 analog of IFN-y, resulted in the expresssion of a single 20 predominant species of polypeptide of the appropriate length.

The following examples 7 and 8 relate to procedures of the invention for generating a manufactured gene specifying the F subtype of human leukocyte interferon ("LeuIFN-F" or "IFN-aF") and polypeptide analogs thereof.

#### EXAMPLE 7

The amino acid sequences for the human leukocyte interferon of the F subtype has been deduced
by way of sequencing of cDNA clones. See, e.g., Goedell,
et al., Nature, 200, pp. 20-26 (1981). The general
procedures of prior Examples 1, 2 and 3 were employed
in the design and assembly of a manufactured DNA sequence
for use in microbial expression of IFN-αF in E. coli
by means of a pBR322-derived expression vector. A



general plan for the construction of three "major" subunit DNA sequences (LeuIFN-F I, LeuIFN-F II and LeuIFN-F III) and one "minor" subunit DNA sequence (LeuIFN-F IV) was evolved and is shown in Table VII below.



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TABLE VII

LeulFN-F IV

BstE II CTAGAGA ATG TGT GAT TTA CCT CAA ACT CAT TCT CTT G
TCT TAC ACA CTA AAT GGA GTT TGA GTA AGA GAA CCATTG -1 1 Met Cys Asp Leu Pro Gln Thr His Ser Leu XbaI

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FCORI 10 GLY ASIN ARG ARG ARG ARG CRC CRC GCT CTG ATT CTG CTG GCA CAG ATG GGT CGT ATT TCC CCG TTT  ABTTCATTA GCT AAC CGT CGC GCT CTG ATT CTG CTG GCA CAG ATG GGT CGT ATT TCC CCG TTT  GTAAT CCA TTG GCA GCG CGA GAC TAA GAC GAC GCT AAC GGC AAA  Bate II  Set Cys Leu Lys Asp Arg His Asp Phe GLY Phe Pro Gln Glu Glu Phe Asp GLY Asg GTT CCG AAC GAC TTC GGC TTT CCG CAA GAA GAC TTT CTG GCA CTG GTG TTT CTG GCA AAG GCG TTT CTG GCA TTT CTG GCA AAG GCG TTT CTG AAG GTT CTT CTC AAG CTA CTG TTT CTG GCA AAG GCG TTT CTG GCA AAG GCG GTT CTT CTC AAG CTG TTT CTG GCA AAG GCG GTT TTT CTG GCA AAG GCG CTT TAC AAG GTT CTG AAA GAC TTT CCG GAA ATG ATC CAG GTG CTT TAC TAG GTT GTG TTT TCC GTA AAG GTT CTA GAG CTT TAC TAG GTT TAC TAG GTT GTG TTT TCC GTC TTT CGA AAA GGC CTT TAG AAG GTT CTG AAA ATG TTT TCC GTC TTT CGA GTT TTT CTG GTT TTT CTG GTT TTT TTT CTG GTT TTT T	4.			he	TTC	Ser	250     260	
Arg Arg Ala Leu Ile Leu Leu Ala Gln Met Gly Arg Ile Ser Processor CGC CGA GAC TAA GAC GCA CAG ATG GGT CGT ATT TCC CGT GCA GCA TAA AGG GCA CGA GAC TAA GAC GAC CGA GTC TAC CCA GCA TAA AGG GGT CGT GTC TAC CCA GCA TAA AGG GGT CGT GTC TAC CCA GCA TAA AGG GGT CGT CTC CAA GAA GAA GAA GAA GAA GAA GAA GAA GA	Phe	TTT		т. H			FA	
Arg Arg Ala Leu Ile Leu Leu Ala Gln Met Gly Arg Ile Ser CGT CGT CGT CGT CGT CGT ATT TCC CGT CGT CGT CGT CGT ATT TCC CGT CGT CGT CGT CGT CGT ATT A GG CGT CGT CGT CGT CGT CGT CGT CGT CGT	ro	ပ္ပပ္တဲ့			25			
Arg Arg Ala Leu Ile Leu Leu Ala Gln Met Gly Arg Ile  39  CGT CGC GCT CTG ATT CTG CTG GCA CAG ATG GGT CGT ATT  I GCA GCG CGA GAC TAA GAC GAC CGT GTC TAC CCA GCA TAA  Asp Arg His Asp Phe Gly Phe Pro Gln Glu Glu Phe Asp Gl  33  34  Ala Ile Ser Val Leu His Glu Met Ile Gln Gln Thr Phe Asp GCT ATC GT TAC GAT GCG GTT TAG GAT GTC TAG AAG TTC GAT GTA GAT TAG ATC TTC GTA GAT GTG TAG GTG TTC TAG AAG TTC TAG AGG TTC TAG TAG ATC TTC TAG AAG TTC TAG AAG TTC TAG AGG TT TAG TAG GTT TAG TAG GTT TAG TAG	H			Asr	AAC	Lei	GAC	
Asp Arg Ala Leu Ile Leu Leu Ala Gln Met Gly Arg Il  CGT CGC GCT CTG ATT CTG CTG GCA CAG ATG GGT CGT AT  GCA GCG CGA GAC TAA GAC GAC CGT GTC TAC CCA GCA TA  Asp Arg His Asp Phe Gly Phe Pro Gln Glu Glu Phe Asp  33  34  Ala Ile Ser Val Leu His Glu Met Ile Gln Gln Thr Phe  GCA ATC TCT GTA CTG CAC GAA ATG ATC CAA CAG ATC  GCA ATC TCT GTA CTG CAC GAA ATG ATC CAA CAG ATC  GGT TAG AGA CTT CAC GAA ATG ATC CAA CAG ACC  GTA ATC TCT GTA CTG CAC GAA ATG ATC CAA CAG AGA  SGT TAG AGA CAT GAC CTT TAC TAG GTT GTC TGG AAG  SER Ala Thr Trp Glu Gln  Hind III	a X	T A		Glγ	ည္မွ	Asn	AAC	
Asp Arg Ala Leu Ile Leu Leu Ala Gln Met G  CGT CGC CGA GAC TAA GAC GAC CGT GTC TAC C  Asp Arg His Asp Phe Gly Phe Pro Gln Glu Glu  Asp Arg CAC GAC Trc GGC Trr CG CAA GAA GAG  CTG GCA GTG CTG AAA GGC GTr CTT  Ala Ile Ser Val Leu His Glu Met Ile Gln Gln  GCA ATC TCT GTA CTG CAC GAA ATG ATC CAA CAG  GCA TAG AG CAT GAC GTT TAC TAG GTT GTC  SGT TAG AGA CAT GAC GTG CTT TAC TAG GTT GTC  Ala Thr Trp Glu Gln  Hind III	11	AT		Q,	AT	he	TC AG	
Asp Arg Ala Leu Ile Leu Leu Ala Gln Met G  CGT CGC CGA GAC TAA GAC GAC CGT GTC TAC C  Asp Arg His Asp Phe Gly Phe Pro Gln Glu Glu  Asp Arg CAC GAC Trc GGC Trr CG CAA GAA GAG  CTG GCA GTG CTG AAA GGC GTr CTT  Ala Ile Ser Val Leu His Glu Met Ile Gln Gln  GCA ATC TCT GTA CTG CAC GAA ATG ATC CAA CAG  GCA TAG AG CAT GAC GTT TAC TAG GTT GTC  SGT TAG AGA CAT GAC GTG CTT TAC TAG GTT GTC  Ala Thr Trp Glu Gln  Hind III	Arg	CGT		e A	ည္သ	r F	-0.00	
Arg Arg Ala Leu Ile Leu Leu  CCGT CGC GCT CTG ATT CTG CTG  Asp Arg His Asp Phe Gly Phe Pr  Asp Arg His Asp Phe Gly Phe Pr  CCT CAC GAC TTC GGC TTT CC  CTG GCA GTG CTG AAG CGG AAA GG  Ala Ile Ser Val Leu His Glu Me  CCA ATC TCT GTA CTG CAC GAA AT  CGT TAG AGA CAT GAC GTG CTT TF  Ser Ala Thr Trp Glu Gln  Hir	31.9	CA		r P	2 E	Ē	T AC	
Arg Arg Ala Leu Ile Leu Leu  CCGT CGC GCT CTG ATT CTG CTG  Asp Arg His Asp Phe Gly Phe Pr  Asp Arg His Asp Phe Gly Phe Pr  CCT CAC GAC TTC GGC TTT CC  CTG GCA GTG CTG AAG CGG AAA GG  Ala Ile Ser Val Leu His Glu Me  CCA ATC TCT GTA CTG CAC GAA AT  CGT TAG AGA CAT GAC GTG CTT TF  Ser Ala Thr Trp Glu Gln  Hir	e t	37— rg o		GII	GAC	<b>G</b> 1.	GAC	
Arg Arg Ala Leu Ile Leu Leu  CCGT CGC GCT CTG ATT CTG CTG  Asp Arg His Asp Phe Gly Phe Pr  Asp Arg His Asp Phe Gly Phe Pr  CCT CAC GAC TTC GGC TTT CC  CTG GCA GTG CTG AAG CGG AAA GG  Ala Ile Ser Val Leu His Glu Me  CCA ATC TCT GTA CTG CAC GAA AT  CGT TAG AGA CAT GAC GTG CTT TF  Ser Ala Thr Trp Glu Gln  Hir	ğ.	CA		<b>G1</b> u	GAA CTT	Gln	CAA	
Arg Arg Ala Leu Ile Leu Leu  CCGT CGC GCT CTG ATT CTG CTG  Asp Arg His Asp Phe Gly Phe Pr  Asp Arg His Asp Phe Gly Phe Pr  CCT CAC GAC TTC GGC TTT CC  CTG GCA GTG CTG AAG CGG AAA GG  Ala Ile Ser Val Leu His Glu Me  CCA ATC TCT GTA CTG CAC GAA AT  CGT TAG AGA CAT GAC GTG CTT TF  Ser Ala Thr Trp Glu Gln  Hir	2 G1	CA		10 11 n	AT.	.]e	YTC 'AG	II
Arg Arg Ala Leu Ile Leu Le  39  C CGT CGC GCT CTG ATT CTG CT  Asp Arg His Asp Phe Gly Phe  3AC CGT CAC GAC TTC GGC TTT  CTG GCA GTG CTG AAG CCG AAA  Ala Ile Ser Val Leu His Glu  ACT TCT GTA CTG CAC GAA  SGT TAG AGA CAT GAC GTG CTT  36  36  37  38  39  39  30  30  30  30  31  31  32  34  35  36  36  37  37  38  38  38  38  38  38  38  38	Ala	GCA		6.0	200		30 TG	
Arg Arg Ala Leu Ile Leu  39 C CGT CGC CGA GAC TAA GAC  Asp Arg His Asp Phe Gly Ph  3AC CGT CAC GAC TTC GGC TT  CTG GCA GTG CTG AAG CCG AA  Ala Ile Ser Val Leu His Gl  3CA ATC TCT GTA CTG CAC GA  3CA ATC TCT GTA CTG CTG CTG  3CA ATC TCT GTA CTG CAC GA  3CA ATC TCT GTA CTG CAC GA  3CA ATC TCT GTA CTG CTG CTG  TG  3CA ATC TCT GTA CTG CTG CTG CTG  3CA ATC TCT GTA CTG	Leu	CTG SAC				79 N	-29-	Hill
ARG ARG ALG LEU  39 C CGT CGC GCA GAC  ASP ARG HIS ASP P  CTG GCA GTG CTG A  SEC ATC TCT GTA C  CGT TAG AGA CAT G  SEC ALG TAG AGA CAT G	e n	TG G		Ph				
ARG ARG ALG LEU  39 C CGT CGC GCA GAC  ASP ARG HIS ASP P  CTG GCA GTG CTG A  SEC ATC TCT GTA C  CGT TAG AGA CAT G  SEC ALG TAG AGA CAT G	์ นั	T C		Gly	885	His	CAC	79 Gln
ASP AR GCA AIA IJ	ĮĮ.	AT		he	T. T. C.	nen		31 u
ASP AR GCA AIA IJ	Leu	CTG		g E	25	I I	AT G	rp 6
ASP AR GCA AIA IJ	Ala	SCT		e A	0 D		F G	H
ASP AR GCA AIA IJ		ည္မွ	T	His				Th
	g A	–39. T C(		Arg	CGT	Ile	ATC	Ala
AATTCATTA GGT AAC GTAAT CCA TTG GTAAT CCA TTG 30 Ser Cys Leu Lys A 35 AGC TGC CTG AAA G TCG ACG GAC TTT C 36 Gln Lys Ala Gln A 31 CAG AAA GCT CAG G GTC TTT CGA GTC 70 Thr Lys Asp Ser 8		၁၁		dsı	AC TG	- Pa	SCA SGT	er Ser
AATTCATTA GGT GTAAT CCA BELF  Ser Cys Leu Ly TCG ACG GAC TY TTG ANA GCT CY GTC TTT CGA GG Thr Lys Asp Se	Asn	AAC	II		14 A 2	, u	25 C C	1 H
AATTCATTA CGATAT CGAGAT CGAGAAT CGAGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGG	10 31y	GT	3stE	Ž.	A FIE	; G	31 4 C	, S
Ser Cys AGC TGC TCG ACG TCG ACG Gln Lys CAG AAA GTC TTT	U		щ	30 Let	GAC	Ala	1200	Asj
ECOR Ser Ser AGC TCG Gln Gln Thr	Ħ	CAT		Cys	TGC	50 Lys	AAA	Lys
	EcoR	AATT		Ser	AGC	Gln	CAG	70 Thr



ACT AAA GAC AGC TCT GCT ACC TGG GAA CAA TGA TTT CTG TCG AGA CGA TGG ACC CTT GTT TCGA

LeuIFN-F II

80 Ser Leu
AATTCAGGCA AGC TTG CTG GAG AAG TTC TCC ACT GTCCGT TCG AAC GAC CTC TTC AAG AGG TGA
Hind III
Leu Asn Asp Met Glu Ala Cys Val Ile Gln Glu Val Gly Val Glu Glu
AAC GAC ATG GAA GCA TTG CTG TAC CTT CGT
-20-

SalI	
113 Met Asn	
Leu	
110 Pro	
Thr	

ACT CCG CTG ATG AAC G
TGA GGC GAC TAC TTG CAGCT



LeuIFN-F I

Leu	1 2 2	I	Phe		TTC		£
	C CTG		Ser		TCT	ģ	I Sal I (end
130 Tyr	TA		150 Arg	n	CA		н
Leu	chg		t t		00		I Sal
hr	CT		Me		TA T		BamHI S
Le J	IC A		116		ATT		щ
ნ. ∺	T A	-71	G1 u		GAA CTT		Stp
n Ar	TAC TTC CAG CGT ATC ACT CTG TAC ATG AGG GTC GCA TAG TGA GAC ATG	•	Trp Glu Val Val Arg Ala Glu Ile Met		CT.	ŀ	166 Arg Leu Arg Arg Lys Glu Stp
Glr	CAC		rg 7		၁ ၁၅		66 1u s
Phe 	TTC		1. A		E A		B G
λ	AC TG		. Va		<b>15</b> 53		LY
rs 7	, 5 5 -		Val		GTA	) P	Arg
a. Ţ	A H		Glu		GAA		Arg
LY	AA		rp (	1,1	. g g		en
ı. Val	GTT				TA		p I
\la	SGA		140 3 Ala		88		Ar Ar
ne 1	rig G		Суs		A C		160 G1u
a Ţ	ATT CTG GCA GTT AAA AAG		Pro		၁၅၅		Gln
Ser Ile Leu Ala Val Lys Lys Tyr Phe Gln Arg Ile Thr Leu	T AT	- <b>1</b>	Ser Pro		rcr AGA		160 Ile Phe Gln Glu
Se	TCT		χr		TA		le
Asp	GAC	H	ស្		A E		ž H
val Val	TC	Sal	ų.		ZE.	<u> </u>	i v
	) 26 26	01	Lys		AAC TTC		Ser
님	cagec dre gae gree cag erg		Glu	ļ	GAA		Leu
Ecori	AATTCAGGC GTC GAC GTCCG CAG CTG	·	e Thr Glu Lys Lys Tyr	-6	ACC GAA AAG AAA TAT TCT CCG TGC GCT TGG GAA GTA GTT CGC GCT GAA ATT ATG CGT TCT TTC TTC TTT TTA AGA GGC ACG CGA ACC CTT CAT CAA GCG CGA CTT TAA TAA GAA GCA AGT AAG		Ser Leu Ser Lys
• •	_ `			- 4		•	

TCT CTG AGC AAA ATC TTC CAG GAG CGT CTG CGC CGT AAA GAA TAA TAG GATCC AGA GAC TCT TTT TAG AAG GTC CTC GCA GAC GCG GCA TTT CTT ATT ATC CTAG



As in the case of the gene manufacture strategy set out in Table IV, the strategy of Table VII involves use of bacterial preference codons wherever it is not inconsistent with deoxyribonucleotide segment 5 constructions. Construction of an expression vector with the subunits was similar to that involved with the IFNy-specifying gene, with minor differences in restriction enzymes employed. Subunit I is ligated into pBR322 cut with EcoRI and SalI. (Note that the 10 subunit terminal portion includes a single stranded SalI "sticky end" but, upon complementation, a SalI recognition site is not reconstituted. A full BamHI recognition site remains, however, allowing for subsequent excision of the subunit.) This first inter-15 mediate plasmid is amplified and subunit II is inserted into the amplified plasmid after again cutting with EcoRI and SalI. The second intermediate plasmid thus formed is amplified and subunit III is inserted into the amplified plasmid cut with EcoRI and HindIII. 20 third intermediate plasmid thus formed is amplified. Subunit IV is ligated to an EcoRI and XbaI fragment isolated from pINTy-TXb4 of Example 4 and this ligation product (having EcoRI and BstEII sticky ends) is then inserted into the amplified third intermediate plasmid 25 cut with EcoRI and BstEII to yield the final expression vector.

The isolated product of trp promoter/operator controlled <u>E.coli</u> expression of the manufactured DNA sequence of Table VII as inserted into the final expression vector was designated IFN- $\alpha F_1$ .

# EXAMPLE 8

As discussed <u>infra</u> with respect to consensus leukocyte interferon, those human leukocyte interferon



subtypes having a threonine residue at position 14 and a methionine residue at position 16 are reputed to display greater antiviral activity than those subtypes possessing Ala<sup>14</sup> and Ile<sup>16</sup> residues. An analog of human leukocyte interferon subtype F was therefore manufactured by means of microbial expression of a DNA sequence of Example 7 which had been altered to specify threonine and methionine as residues 14 and 16, respectively. More specifically, [Thr $^{14}$ , Met $^{16}$ ] IFN- $\alpha$ F, designated IFN-oF2, was expressed in E.coli upon transformation with a vector of Example 7 which had been cut with SalI and HindIII and into which a modified subunit II (of Table VII) was inserted. The specific modifications of subunit II involved assembly with seg-15 ment 39 altered to replace the alanine-specifying codon, GCT, with a threonine-specifying ACT codon and replace the isoleucine-specifying codon, ATT, with an ATG codon. Corresponding changes in complementary bases were made in section 40 of subunit LeuIFN-FII. The following Examples 9 and 10 relate to 20 practice of the invention in the microbial synthesis

of consensus human leukocyte interferon polypeptides which can be designated as analogs of human leukocyte interferon subtype F.

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#### EXAMPLE 9

"Consensus human leukocyte interferon" ("IFN-Con," "LeuIFN-Con") as employed herein shall mean a non-30 naturally-occurring polypeptide which predominantly includes those amino acid residues which are common to all naturally-occurring human leukocyte interferon subtype sequences and which includes, at one or more of those positions wherein there is no amino acid common to all subtypes, an amino acid which predomi-



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nantly occurs at that position and in no event includes any amino acid residue which is not extant in that position in at least one naturally-occurring subtype. (For purposes of this definition, subtype A is positionally aligned with other subtypes and thus reveals a "missing" amino acid at position 44.) As so defined, a consensus human leukocyte interferon will ordinarily include all known common amino acid residues of all subtypes. will be understood that the state of knowledge con-10 cerning naturally-occurring subtype sequences is continuously developing. New subtypes may be discovered which may destroy the "commonality" of a particular residue at a particular position. Polypeptides whose structures are predicted on the basis of a later-amended determination of commonality at one or more positions would 15 remain within the definition because they would nonetheless predominantly include common amino acids and because those amino acids no longer held to be common would nonetheless quite likely represent the predominant amino acid at the given positions. Failure of 20 a polypeptide to include either a common or predominant amino acid at any given position would not remove the molecule from the definition so long as the residue at the position occurred in at least one subtype. Polypeptides lacking one or more internal or terminal residues of consensus human leukocyte interferon or including internal or terminal residues having no counterpart in any subtype would be considered analogs of human consensus leukocyte interferon.

Published predicted amino acid sequences for eight cDNA-derived human leukocyte interferon subtypes were analyzed in the context of the identities of amino acids within the sequence of 166 residues. See, generally, Goedell, et al., Nature, 290, pp. 20-26 (1981) comparing LeIFN-A through LeIFN-H and noting that only 79 amino acids appear in identical positions in all



eight interferon forms and 99 amino acids appear in identical positions if the E subtype (deduced from a cDNA pseudogene) was ignored. Each of the remaining positions was analyzed for the relative frequency of occurrence of a given amino acid and, where a given amino acid appeared at the same position in at least five of the eight forms, it was designated as the predominant amino acid for that position. A "consensus" polypeptide sequence of 166 amino acids was plotted out and compared back to the eight individual sequences, resulting in the determination that LeIFN-F required few modifications from its "naturally-occurring" form to comply with the consensus sequence.

A program for construction of a manufactured

15 IFN-Con DNA sequence was developed and is set out
below in Table VIII. In the table, an asterisk designates the variations in IFN-αF needed to develop
LeIFN-Con<sub>1</sub>, i.e., to develop the [Arg<sup>22</sup>, Ala<sup>76</sup>, Asp<sup>78</sup>,
Glu<sup>79</sup>, Tyr<sup>86</sup>, Tyr<sup>90</sup>, Leu<sup>96</sup>, Thr<sup>156</sup>, Asn<sup>157</sup>, Leu<sup>158</sup>]

20 analog of IFN-αF. The illustrated top strand sequence includes, wherever possible, codons noted to the subject of preferential expression in E. coli. The sequence also includes bases providing recognition sites for Sal,
HindIII, and BstE2 at positions intermediate the sequence and for XBaI and BamHI at its ends. The latter sites are selected for use in incorporation of the sequence in a pBR322 vector, as was the case with the sequence developed for IFN-αF and its analogs.

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TAG

# TABLE VIII

	-1	7				:		•	•	•	10			
	Mot.	-Cvc	-Asp- GAT	-Leu- TTA	-Pro- CCT	-Gln CAA	-Thr ACT	-His CAT	-Ser TCT	-Leu- CTT	-G1 y-	-Asn- AAC	-Arg- CGT	-Arg- CGC
5	Ala GCT	-Leu CTG	-Ile ATT	-Leu- CTG	-Leu- CTG	-Ala GCA	20 -Gln CAG	-Met ATG	* -Arg- CGT	-Arg CGT	-Ile- ATT	-Ser TCC	-Pro- CCG	-Phe- TTT
	Ser-	-Cys- TGC	30 -Leu- CTG	-Lys- AAA	-Asp GAC	-Arg CGT	-His- CAC	-Asp- GAC	-Phe- TTC	-Gly- GGC	-Phe-	-Pro- CCG	40 -Gln- CAA	-Glu- GAA
10	Glu- GAG	-Phe TTC	-Asp- GAT	-Gly- GGC	-Asn- AAC	-Gln- CAA	-Phe TTC	-Gln- CAG	50 -Lys- AAA	-Ala- GCT	-Gln- CAG	-Ala- GCA	-Ile- ATC	-Ser- TCT
15	Val- GTA	-Leu- CTG	-His- CAC	-Glu- GAA	60 -Met- ATG	-Ile- ATC	-Gln- CAA	-Gln- CAG	-Thr - ACC	-Phe- TTC	-Asn- AAC	-Leu- CTG	-Phe- TTT	-Ser- TCC
15	70 Thr- ACT	-Lys- AAA	-Asp- GAC	-Ser- AGC	-Ser- TCT	-Ala- GCT	* -Ala- GCT	-Trp- TGG	* -Asp- GAC	* -Glu- GAA	80 Ser- AGC	-Leu- TTG	-Leu- CTG	-Glu- GAG
20	Lys-	-Phe- TTC	* -Tyr- TAC	-Thr - ACT	-Glu- GAA	-Leu- CTG	*90 -Tyr- TAT	-Gln- CAG	-Gln- CAG	-Leu- CTG	-Asn- AAC	-Asp- GAC	* -Leu- CTG	-Glu- GAA
	Ala- GCA	-Cys- TGC	100 -Val- GTA	-Ile- ATC	-Gln- CAG	-Glu- GAA	-Val- GTT	-Gly- GGT	-Val- GTA	-Glu- GAA	-Glu- GAG	-Thr - ACT	110 Pro- CCG	-Leu- CTG
25	Met- ATG	-Asn- AAC	-Val- GTC	-Asp- GAC	-Ser- TCT	-Ile- ATT	-Leu- C <b>T</b> G	-Ala- GCA	120 -Val- GTT	-Lys- AAA	-Lys- AAG	-Tyr- TAC	-Phe- TTC	-Gln- CAG
	Arg- CGT	-Ile- ATC	-Thr - ACT	-Leu- CTG	130 Tyr- TAC	-Leu- CTG	-Thr- ACC	-Glu- GAA	-Lys- AAG	-Lys- AAA	-Tyr- TAT	-Ser- TCT	-Pro- CCG	-Cys- TGC
30	140 Ala- GCT	-Trp- TGG	-Glu- GAA	-Val- GTA	-Val- GTT	-Arg- CGC	-Ala- GCT	-Glu- GAA	-Ile- ATT	-Met- ATG	150 Arg- CGT	-Ser- TCT	TTC	
	Leu- CTG	-Ser- TCT	* -Thr- ACT	* -Asn- AAC	* -Leu- CTG	-Gln- CAG	160 -Glu- GAG	-Arg- CGT	-Leu- CTG	-Arg- CGC	Arg- CGT	-Lys- AAA	-Glu	Stop TAA
35	Stop	Þ												•



- 64 -

Table IX below sets out the specific double stranded DNA sequence for preparation 4 subunit DNA sequences for use in manufacture of IFN-Con<sub>1</sub>. Subunit LeuIFN-Con IV is a duplicate of LeuIFN-F IV of Table VIII. Segments of subunits which differ from those employed to construct the IFN-αF gene are designated with a "prime" (e.g., 37' and 38' are altered forms of sections 37 and 38 needed to provide arginine rather than glycine at position 22).

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TABLE IX

LeuIFN Con IV

BstE II CTAGAGA ATG TGT GAT TTA CCT CAA ACT CAT TCT CTT G
TCT TAC ACA CTA AAT GGA GTT TGA GTA AGA GAA CCATTG -1 1 2 3 4 5 6 7 8 9 Met Cys Asp Leu Pro Gln Thr His Ser Leu xbai

> BUREAU OMPI WIPO WIPO WIPO

- 66 -

ECORI	1 1	10 Gly Asn GGT AAC	1			Ala Leu GCT CTG	i i	Ile Leu	Leu Le	Leu Ala		20 In Met	let Arg	Arg Arg				o Phe	υ 1 E-1 i
Cys	C I	Ly Ly		sp Arg	<i>)</i>	40————————————————————————————————————	gp Phe	≰   5			40 Gln G	Glu G	٦ او	38	Asp G	Gly As	Asn G	Gln P	A Phe
TGC	C CTG	35- AAA TTT 36-	GAC	CGT	CAC	GAC	Trc	၁၉၅	TTT	၁၁၁	CAA		GAG	TTC	GAT	၁၁၁	AAC	CAA	TTC
50 Lys	<<	d Gln	Ala	Ile	Ser	Val	Leu	His	<b>61</b> u	60 Met	The	Gln	Gln Thr		Phe	Asn	Leu	Phe	Ser
AAA	88	-31 CT CAG SA GTC	GCA	ATC		TCT GTA	GAC	CAC	GAA	A ATG	ATC	CAA	CAG	ACC	TTC	AAC	CTG	TTT	TCC
Lys	Lys Asp	Ser	Ser	Ala		79 Ala Trp Asp Glu	Asp	79 Glu	H	Hind I	III								
AAA TTT	A GAC r crg	AGC TCG	TCT	GCT	GCT CGA	HK	TGG GAC ACC CTG	GAA	TCGA	۳٦									



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LeuIFN Con II

In	CAG		Glu	GAG
e E	CAG CAG GTC GTC		Glu	GAA
ි 	6.0		1.	AT TA
90 Tyr	TAT		Š	50
ren	CTG		G13	595
lu 1	AG C	; 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	Val	GTT CAA 18—
. G	ACC GAG CTG TAT TGG CTC GAC ATA	·	Ile Gln Glu Val Gly Val Glu	GAA
를 c	0 0 E		1, n	CAG GTC
Тy	TA		0	ပြုပ္ချ
he	FTC	<u> </u>  -	II	TA
80 Ser Leu Leu Glu Lys Phe Tyr Thr Glu Leu Tyr Gln Gln	AG 1		100 Val	CTG GAA GCA TGC GTA ATC CAG GAA GTT GGT GTA GAC CTT CGT ACG CAT TAG GTC CTT CAA CCA CAT 20'
I u	AG A		Cγs	TGC
<u>ල</u>	90		ø	ST
rèn	CTC		Z.	88
ne n	PAC PAC	-774 11	Glu	GAA CTT
80 er I	ည္ပည္	d III	Leu	CTG
, w	A INC.	Hind	Asp	GAC CTG
	AATTCAGGCA AGC TTG CTG GAG AAG TTC TAC ACC GTCCGT TCG AAC GAC CTC TTC AAG ATG TGG		Leu Asn Asp Leu Glu Ala Cys	AAC GAC (TITG CTG (
ECORI	] [ 5년 - 6	1	<b>₹</b>	CTG A
E E	[\\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\		Le	1581

ACT CCG CTG ATG AAC G
TGA GGC GAC TAC TTG CAGCT 110 113 Thr Pro Leu Met Asn

Sal I



LeuIFN Con I

3	1001 .	Phe	TTC	_	
Leu	TAC CTG ATG GAC	Ser	TCT AGA	Sal I (end)	
130 TYr	TAC	$\sim$ $\sim$	CGT T	н	
Leu	cre rac gac arg	t P	ပ္ ပ	Sal	ည်
rhr.	ACT	w We	r AT	BamHI	<u>ည်ခြီ</u>
le :	ATC PTAG 12	Ţ	AT		CT
I 6	TA A	G1u	GAA	Stp	TAG
n Aı	99	Ala	GGA.	Stp	TAA TAG GATCC ATT ATC CTAGGAGCT
G1	TTC CAG CGT ATC AAG GTC GCA TAG	Arg	ပ္ပစ္တပ္တ	166 Glu	SAA
Lys Lys Tyr Phe Gln Arg Ile Thr Leu	AAA AAG TAC TTC CAG CGT ATC ACT CTG TTT TTC ATG AAG GTC GCA TAG TGA GAC 12	Trp Glu Val Val Arg Ala Glu Ile Met	TT (	ე გ	AA (TT
Tyr	AAA AAG TAC TTT TTC ATG	а1 V	TA G	160 Glu Arg Leu Arg Arg Lys	GA T
Lys	AAG	n A	TA G	ğ ğ	ပ္ ဗ
Ĺys	AAA	ь С	ซีบี	u Ar	- ၁ ၁ ၁ ၁
120 Val	GTT	Trp	A A C	Le	CT
la V	GCA G	140 Ala	GGA	Arg	CGI
u A	CTG GCA GAC CGT	Cys	TGC ACG	160 Glu	GAG CTC
e Le	T CTG	Pro Cys	၁၁၁	GIn	CAG
Ser Ile Leu Ala	TCT ATT AGA TAA		TCT CCG TGC GCT TGG GAA GTA GTT CGC GCT GAA ATT ATG CGT TCT TTC AGA GGC ACG CGA ACC CTT CAT CAA GCG CGA CTT TAA TAC GCA AGA AAG 6	Leu	AAC CTG CAG GAG CGT CTG CGC CGT AAA GAA TAA TAG GATCC TTG GAC GTG GCA TTT CTT ATT ATC CTAGS
	-13 C TCT G AGA	Tyr Ser	TAT A	san I	AC. C
114 Val Asp			1 - 1	A.	
114 Val	CAGGC GTC GAC GTCCG CAG CTC Sal I	Glu Lys Lys	C P	된	AGC ACT TCG TGA
	ပ္တစ္ပ	ı Ly	A AA	s Se	AG TC
RI	AATTCAGGC GTC GAG GTCCG CAG CT Sal I	Glu	ACC GAA AAG AAA TGG CTT TTC TTT	Ser Leu Ser Thr	rcr cre agc acr aga gac rcg rga
ECORI	AAT	Thr	ACC	Ser	TCT



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The four subunits of Table IX were sequentially inserted into an expression vector according to the procedure of Example 7 to yield a vector having the coding region of Table VIII under control of a trp promoter/ operator. The product of expression of this vector in E.coli was designated IFN-Con1. It will be noted that this polypeptide includes all common residues indicated in Goedell, et al., supra, and, with the exception of Ser 80, Glu 83, Val 114, and Lys 121, included the predominant amino acid indicated by analysis of the reference's summary of sequences. The four above-noted residues were retained from the native IFN-αF sequence to facilitate construction of subunits and assembly of subunits into an expression vector. (Note, e.g., serine was retained at position 80 to allow for construction of a HindIII site.)

Since publication of the Goedell, et al. summary of IFN- $\alpha$  subtypes, a number of additional subtypes have been ascertained. Figure 2 sets out in tabular form the deduced sequences of the 13 presently known subtypes (exclusive of those revealed by five known cDNA pseudogenes) with designations of the same IFN-α subtypes from different laboratories indicated parenthetically (e.g., IFN- $\alpha$ 6 and IFN- $\alpha$ K). See, e.g., Goedell, et al., supra; Stebbing, et al., in: Recombinant DNA Products, Insulin, Interferons and Growth Hormones (A. Bollon, ed.), CRC Press (1983); and Weissman, et al., U.C.L.A. Symp.Mol.Cell Biol., 25, pp. 295-326 (1982). Positions where there is no common amino acid are shown in bold face. IFN- $\alpha$  subtypes are roughly grouped on the basis of amino acid residues. In seven positions (14, 16, 71, 78, 79, 83, and 160) the various subtypes show just two alternative amino acids, allowing classification of the subtypes into two subgroups (I and II) based on which of the seven positions are occupied by the same amino acid residues.



Three IFN-α subtypes (H, F and B) cannot be classified as Group I or Group II and, in terms of distinguishing positions, they appear to be natural hybrids of both group subtypes. It has been reported that IFN-α subtypes of the Group I type display relatively high antiviral activity while those of Group II display relatively high antitutely high antitutely high antitutely.

line of the Figure. It is noteworthy that certain residues of IFN-Con<sub>1</sub> (e.g., serine at position 8) which were determined to be "common" on the basis of the Goedell, et al., sequences are now seen to be "predominant". Further, certain of the IFN-Con<sub>1</sub> residues determined to be predominant on the basis of the reference (Arg<sup>22</sup>, Asp<sup>78</sup>, Glu<sup>79</sup>, and Tyr<sup>86</sup>) are no longer so on the basis of updated information, while certain heretofore non-predominant others (Ser<sup>80</sup> and Glu<sup>83</sup>) now can be determined to be predominant.

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# EXAMPLE 10

A human consensus leukocyte interferon which differed from IFN-Con<sub>1</sub> in terms of the identity of amino acid residues at positions 14 and 16 was prepared by modification of the DNA sequence coding for IFN-Con<sub>1</sub>. More specifically, the expression vector for IFN-Con<sub>1</sub> was treated with BstEII and Hind III to delete subunit LeuIFN Con III. A modified subunit was inserted wherein the alanine-specifying codon, GCT, of sections 39 and 40 was altered to a threonine-specifying codon, ACT, and the isoleucine codon, CTG, was changed to ATG. The product of expression of the modified manufactured gene, [Thr<sup>14</sup>, Met<sup>16</sup>, Arg<sup>22</sup>, Ala<sup>76</sup>, Asp<sup>78</sup>, Glu<sup>79</sup>, Tyr<sup>86</sup>, Tyr<sup>90</sup>, Leu<sup>96</sup>, Thr<sup>156</sup>, Asn<sup>157</sup>, Leu<sup>158</sup>]IFN-αF, was designated IFN-Con<sub>2</sub>.



Presently being constructed is a gene for a consensus human leukocyte interferon polypeptide which will differ from IFN-Con, in terms of the identity of residues at positions 114 and 121. More specifically, the Val 114 and Lys 121 residues which duplicate IFN-of subtype residues but are not predominant amino acids will be changed to the predominant  $Glu^{114}$  and  $Arg^{121}$ residues, respectively. Because the codon change from Val 114 to Arg 114 (e.g., GTC to GAA) will no longer allow for a SalI site at the terminal portion of subunit 10 LeuIFN Con I (of Table IX), subunits I and II will likely need to be constructed as a single subunit. Changing the AAA, lysine, codon of sections 11 and 12 to CTG will allow for the presence of arginine at position 121. The product of microbial expression of the manufactured gene, [Arg<sup>22</sup>, Ala<sup>76</sup>, Asp<sup>78</sup>, Glu<sup>79</sup>, Tyr<sup>86</sup>, Tyr<sup>90</sup>, Leu<sup>96</sup>, Glu<sup>114</sup>, Arg<sup>121</sup>, Thr<sup>156</sup>, Asn<sup>157</sup>, Leu<sup>158</sup>] IFN-aF, will be designated IFN-Con3.

The following example relates to procedures for enhancing levels of expression of exogenous genes in bacterial species, especially, <u>E.coli</u>.

### EXAMPLE 11

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In the course of development of expression vectors in the above examples, the trp promoter/operator DNA sequence was employed which included a ribosome binding site ("RBS") sequence in a position just prior to the initial translation start (Met -1, ATG). An attempt was made to increase levels of expression of the various exogenous genes in <u>E.coli</u> by incorporating DNA sequences duplicative of portions of putative RBS sequences extant in genomic <u>E.coli</u> DNA sequences associated with highly expressed cellular proteins. Ribosome binding site sequences of such protein-coding genes as



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reported in Inokuchi, et al. <u>Nuc.Acids.Res.</u>, <u>10</u>, pp. 6957-6968 (1982), Gold, et al., <u>Ann.Rev.Microbiol.</u>, <u>35</u>, pp. 365-403 (1981) and Alton, et al., <u>Nature</u>, <u>282</u>, pp. 864-869 (1979), were reviewed and the determination was made to employ sequences partially duplicative of those associated with the <u>E.coli</u> proteins OMP-F (outer membrane protein F), CRO and CAM (chloramphenicol transacetylase).

By way of example, to duplicate a portion of the OMP-F RBS sequence the following sequence is inserted prior to the Met-1 codon.

5'-AACCATGAGGGTAATAAATA-3'
3'-TTGGTACTCCCATTATTTAT-5'

In order to incorporate this sequence in a position prior to the protein coding region of, e.g., the manufactured gene coding for IFN-Con<sub>1</sub> or IFN-αF<sub>1</sub>, subunit IV of the expression vector was deleted (by cutting the vector with XbaI and BstEII) and replaced with a modified subunit IV involving altered sections 41A and 42A and the replacement of sections 43 and 44 with new segments RBl and RB2. The construction of the modified sequence is as set out in Table X, below.

25 .

### TABLE X

XbaI

-1 1 2 Met Cys Asp

RB1

30 CTAGAAA CCA TGA GGG TAA TAA ATA ATG TGT GAT
TTT GGT ACT CCC ATT ATT TAT TAC ACA CTA

RB2

3 4 5 6 7 8 9 Leu Pro Gln Thr His Ser Leu BstEII

TTA CCT CAA ACT CAT TCT CTT G
AAT GGA GTT TGA GTA AGA GAA CATG



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Table XI, below, illustrates the entire DNA sequence in the region preceding the protein coding region of the reconstructed gene starting with the HpaI site within the trp promoter/operator (compare subunit IF-4 of Table IV).

# TABLE XI

HpaI

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XbaI

10 AAC TAG TAC GCA AGT TCA CGT AAA AAG GGT ATC TAG AAA CCA TTG ATC ATG CGT TCA AGT GCA TTT TTC CCA TAG ATC TTT GGT

-1 1 2 3 4 5 6 7 Met Cys Asp Leu Pro Gln Thr His

TGA GGG TAA TAA ATA ATG TGT GAT TTA CCT CAA ACT CAT ACT CCC ATT ATT TAC ACA CTA AAT GGA GTT TGA GTA

8 9 BstE II Ser Leu TCT CTT G AGA GAA CATG

Similar procedures were followed to incorporate sequences duplicative of RBS sequences of CRO and CAM genes, resulting in the following sequences immediately preceding the Met-1 codon.

25 1 10 20
CRO: GCATGTACTAAGGAGGTTGT
CGTACATGATTCCTCCAACA

1 10 20

CAM: CAGGAGCTAAGGAAGCTAAA GTCCTCGATTCCTTCGATTT

It will be noted that all the RBS sequence inserts possess substantial homology to Shine-Delgarno sequences, are rich in adenine and include sequences ordinarily providing "stop" codons.



Levels of E.coli expression of IFN-Con, were determined using trp-controlled expression vectors incorporating the three RBS inserts (in addition to the RBS sequence extant in the complete trp promoter/operator): Expression of the desired polypeptide using the OMP-F RBS duplicating sequence was at from 150-300 mg per liter of culture, representing from 10 to 20 percent of total protein. Vectors incorporating the CAM RBS duplicating sequence provided levels of expression which 10 were about one-half that provided by the OMP-F variant. Vectors including the CRO RBS duplicating sequence yielded the desired protein at levels of about one-tenth that of the OMP-F variant.

The following example relates to antiviral 15 activity screening of human leukocyte interferon and polypeptides provided by the preceding examples.

## EXAMPLE 12

Table XII below provides the results of testing of antiviral activity in various cell lines of natural (buffy coat) interferon and isolated, microbiallyexpressed, polypeptides designated IFN-aF1, IFN-aF2, IFN-Con,, and IFN-Con, Viruses used were VSV (vesicular 25 stomatitis virus) and EMCV (encephalomyocarditis virus). Cell lines were from various mammalian sources, including human (WISH, HeLa), bovine (MDBK), mouse (MLV-6), and monkey (Vero). Antiviral activity was determined by an end-point cytopathic effect assay as described in Weck, 30 et al., J.Gen.Virol., 57, pp. 233-237 (1981) and Campbell, et al., Can.J.Microbiol., 21, pp. 1247-1253 (1975). Data shown was normalized for antiviral activity in WISH cells.



# TABLE XII

	Virus	Cell Line	Buffy Coat	IFN- <sup> </sup>	IFN- <sup>αF</sup> 2	Con <sub>1</sub>	IFN- Con <sub>2</sub>	
5	vsv	WISH 5						digally some of the
:	vsv	HeLa	400	100	'*ND*	1997 <b>200</b> %	100	海南 双联 不然 医电影 经
	vsv	MDBK	1600	33	ND	200	300	
	vsv	MLV-6	20	5	ND	3	20	•
	vsv	Vero	10	0.1	ND	10	0.	1
10	EMCV	WISH	100	100	100	100	100	·
	EMCV	HeLa	100	5	ND	33	33	
	EMCV	Vero	100	20	·ND·	1000	10	

<sup>\*</sup>ND - no data presently available.

15 It will be apparent from the above examples that the present invention provides, for the first time, an entire new genus of synthesized, biologically active proteinaceous products which products differ from naturally-occurring forms in terms of the identity

20 and/or location of one or more amino acids and in terms of one or more biological (e.g., antibody reactivity) and pharmacological (e.g., potency or duration of effect) but which substantially retain other such properties.

products of the present invention and/or antibodies thereto may be suitably "tagged", for example
radiolabelled (e.g., with I<sup>125</sup>) conjugated with enzymes
or fluorescently labelled, to provide reagent materials
useful in assays and/or diagnostic test kits, for the
qualitative and/or quantitative determination of the
presence of such products and/or said antibodies in
fluid samples. Such anitbodies may be obtained from
the innoculation of one or more animal species (e.g.,
mice rabbit, goat, human, etc.) or from monoclonal antibody sources. Any of such reagent materials may be used
alone or in combination with a suitable substrate, e.g.,
coated on a glass or plastic particle bead.



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Numerous modifications and variations in the practice of the invention are expected to occur to those skilled in the art upon consideration of the foregoing illustrative examples. Consequently, the invention should be considered as limited only to the extent reflected by the appended claims. ...



#### WHAT IS CLAIMED IS:

- 1. A method for the manufacture of linear, double stranded DNA sequences of a length in excess of about 200 base pairs and coding for expression of a predetermined continuous sequence of amino acids with a selected host microorganism transformed by a selected DNA vector including the sequence, said method comprising:
- 10 (a) preparing two or more different, subunit, linear, double stranded DNA sequences of from about 100 to about 200 base pairs in length,

each different subunit DNA sequence comprising a series of nucleotide base codons coding for a different continuous portion of said predetermined sequence of amino acids to be expressed,

one terminal region of a first of said subunits comprising a portion of a base sequence which
provides a recognition site for cleavage by a first
restriction endonuclease, which recognition site is
entirely present either once or not at all in said
selected assembly vector upon insertion of the subunit
therein,

one terminal region of a second of said
subunits comprising a portion of a base sequence which
provides a recognition site for cleavage by a second
restriction endonuclease other than said first endonuclease, which recognition site is entirely present
once or not at all in said selected assembly vector
upon insertion of the subunit therein,

at least one-half of all remaining terminal regions of subunits comprising a portion of a recognition site for restriction endonuclease cleavage by an endonuclease other than said first and second endonucleases, which recognition site is entirely present once and only once in said selected assembly vector after insertion of all subunits thereinto; and



- (b) serially inserting each of said subunit

  DNA sequences prepared in step (a) into the selected
  assembly vector and effecting the biological amplification of the assembly vector subsequent to each insertion,
  thereby to form a DNA vector including the desired
  DNA sequence coding for the predetermined continuous
  amino acid sequence and wherein the desired DNA sequence
  assembly includes at least one unique recognition
  site for restriction endonuclease cleavage at an intermediate position therein.
- A method according to claim 1 wherein the restriction site for endonuclease cleavage by the restriction endonuclease other than said first
   and second endonucleases is a palindromic six base recognition site and the desired DNA sequence assembled has at least one unique six base recognition site for restriction endonuclease cleavage at an intermediate position therein.

3. A method according to claim 1 wherein at least three different subunit DNA sequences are prepared in step (a) and serially inserted into said selected vector in step (b) and the desired DNA sequence obtained includes at least two unique restriction endonuclease recognition sites at intermediate positions therein.

- 4. A method according to claim 1 wherein
  30 the DNA sequence manufactured comprises an entire
  structural gene coding for a biologically active polypeptide.
- 5. A method according to claim 1 wherein, 35 in the DNA sequence manufactured, the sequence of nucleotide bases includes one or more codons selected,



from among alternative codons specifying the same amino acid, on the basis of preferential expression characteristics of the codon in said selected host microorganism.

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DNA sequence of a length in excess of about 200 base pairs and coding for the expression of a predetermined continuous sequence of amino acids by a selected host microorganism transformed with a selected DNA vector including the sequence, said sequence characterized by having at least one unique palindromic six base recognition site for restriction endonuclease cleavage at an intermediate position therein.

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- 7. A DNA sequence according to claim 6 characterized by having two or more unique restriction endonuclease cleavage sites at intermediate positions therein, at least one of which has a six base palindromic recognition site.
  - 8. A DNA sequence according to claim 6 which comprises an entire structural gene coding for a biologically active polypeptide.

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- 9. A DNA sequence according to claim 8 wherein the gene codes for a human polypeptide.
- 10. A DNA sequence according to claim 8
  30 wherein the gene codes for a polypeptide which differs from a naturally-occurring human polypeptide in terms of the identity and/or location of one or more amino acids.
- 35 11. A DNA sequence according to claim 6 wherein the sequence of nucleotide bases includes



at least one or more codons selected, from among alternative codons specifying the same amino acids, on the basis of preferential expression characteristics of the codon in said selected host microorganism.

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- 12. A polypeptide product of the expression by an organism of a manufactured DNA sequence according to claim 6.
- 13. A biologically functional DNA microorganism transformation vector including a manufactured DNA sequence according to claim 6.
- 14. A vector according to claim 12 which 15 is a circular DNA plasmid.
  - 15. A manufactured gene capable of directing the synthesis in a selected host microorganism of human immune interferon.

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16. A manufactured gene according to claim
15 having at least one unique six base recognition
site for restriction endonuclease cleavage at an intermediate position therein.

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- 17. A manufactured gene according to claim
  15 wherein the base sequence includes one or more
  codons selected, from among alternative codons specifying the same amino acid, on the basis of preferential
  expression characteristics of the codon in a projected
  host microorganism.
- 18. A manufactured gene according to claim
  17 wherein the base sequence includes one or more
  35 codons selected, from among alternative codons specifying the same amino acid, on the basis of preferential
  expression characteristics of the codon in <u>E</u>. <u>coli</u>.



- 19. A manufactured gene according to claim
  15 wherein the base sequence, commencing with the
  5' end of the top strand, is as follows:
- 1 TGT-TAC-TGC-CAG-GAT-CCG-TAC-GTT-AAG-GAA-GCA-GAA-AAC-CTG-
- 15 AAA-AAA-TAC-TTC-AAC-GCA-GGC-CAC-TCC-GAC-GTA-GCT-GAT-AAC-
- 29 GGC-ACC-CTG-TTC-CTG-GGT-ATC-CTA-AAA-AAC-TGG-AAA-GAG-GAA-
- 43 TCC-GAC-CTG-AAG-ATC-ATG-CAG-TCT-CAA-ATT-GTA-AGC-TTC-TAC-
- 57 TTC-AAA-CTG-TTC-AAG-AAC-TTC-AAA-GAC-GAT-CAA-TCC-ATC-CAG-
- 71 AAG-AGC-GTA-GAA-ACT-ATT-AAG-GAG-GAC-ATG-AAC-GTA-AAA-TCC-
- 85 TTT-AAC-AGC-AAC-AAG-AAG-AAA-CGC-GAT-GAC-TTC-GAG-AAA-CTG-
- 94 ACT-AAC-TAC-TCT-GTT-ACA-GAT-CTG-AAC-GTG-CAG-CGT-AAA-GCT-
- 113 ATT-CAC-GAA-CTG-ATC-CAA-GTT-ATG-GCT-GAA-CTG-TCT-CCT-GCG-
- 127 GCA-AAG-ACT-GGC-AAA-CGC-AAG-CGT-AGC-CAG-ATG-CTG-TTT-CAG-
- 141
  [or CGT]-CGT-CGC-CGT-GCT-TCT-CAG.
- 20. A manufactured gene according to claim 19 wherein the base codon for amino acid 41 is GAA and for amino acid 46 is CGC.
  - 21. A biologically functional DNA microorganism transformation vector including a manufactured gene according to claim 15.



- 22. A vector according to claim 21 which is a circular DNA plasmid.
- 23. A process for the production of human 5 immune interferon comprising:

growing, under appropriate nutrient conditions, microorganisms transformed with biologically functional DNA including a manufactured gene according to claim 15, whereby said microorganisms express said gene and produce authentic human immune interferon.

- 24. A process according to claim 23 wherein the microorganisms grown are E. coli microorganisms.
- 15 25. A manufactured gene capable of directing the synthesis in a selected host microorganism of a polypeptide which differs from human immune interferon in terms of the identity and/or location of one or more amino acids.

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26. A manufactured gene according to claim 25 having at least one unique six base recognition site for restriction endonuclease cleavage at an intermediate position therein.

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- 27. A manufactured gene according to claim 25 wherein the base sequence includes one or more codons selected, from among alternative codons specifying the same amino acid, on the basis of preferential expression characteristics of the codon in a projected host microorganism.
- 28. A manufactured gene according to claim
  27 wherein the base sequence includes one or more codons
  35 selected, from among alternative codons specifying the
  same amino acid, on the basis of preferential expression
  characteristics of the codon in <u>E</u>. <u>coli</u>.

- 29. A biologically functional DNA microorganism transformation vector including a manufactured gene according to claim 25.
- 30. A vector according to claim 29 which is a circular DNA plasmid.
- peptide analog of human immune interferon comprising:

  growing, under appropriate nutrient conditions,
  microorganisms transformed with biologically functional
  DNA including a manufactured gene according to claim 25,
  whereby said microorganisms express said gene and produce
  authentic human immune interferon.
- 15 32. A process according to claim 31 wherein the microorganisms grown are  $\underline{E}$ .  $\underline{coli}$  microorganisms.
- 33. A polypeptide product of the process of claim 23.
  - 34. A polypeptide product of the process of claim 31.
- 35. A polypeptide product according to claim 34 selected from the group consisting of:

[Lys<sup>81</sup>]IFNY; [Ser<sup>1</sup>, Ser<sup>3</sup>, Lys<sup>81</sup>]IFNY; [Lys<sup>1</sup>, Lys<sup>2</sup>, Gln<sup>3</sup>, Lys<sup>81</sup>]IFNY; [des-Cys<sup>1</sup>, des-Tyr<sup>2</sup>, des-Cys<sup>3</sup>, Lys<sup>81</sup>]IFNY; [Phe<sup>39</sup>, Lys<sup>81</sup>]IFNY; [Thr<sup>48</sup>, Lys<sup>81</sup>]IFNY; [Lys<sup>81</sup>, Cys<sup>95</sup>]IFNY; [Cys<sup>95</sup>]IFNY; [Cys<sup>95</sup>, Pro<sup>104</sup>]IFNY; [Gln<sup>28</sup>]IFNY; and [Glu<sup>5</sup>]IFNY.

- 36. A manufactured gene capable of directing the synthesis in a selected host microorganism of consensus human leukocyte interferon.
- 5 37. Consensus human leukocyte interferon.

- 38. A consensus human leukocyte interferon according to claim 37 selected from the group consisting of:
- [Arg<sup>22</sup>, Ala<sup>76</sup>, Asp<sup>78</sup>, Glu<sup>79</sup>, Tyr<sup>86</sup>, Tyr<sup>90</sup>,

  Leu<sup>96</sup>, Thr<sup>156</sup>, Asn<sup>157</sup>, Leu<sup>158</sup>]IFN-αF;

  [Thr<sup>14</sup>, Met<sup>16</sup>, Arg<sup>22</sup>, Ala<sup>76</sup>, Asp<sup>78</sup>, Glu<sup>79</sup>,

  Tyr<sup>86</sup>, Tyr<sup>90</sup>, Leu<sup>96</sup>, Thr<sup>156</sup>, Asn<sup>157</sup>, Leu<sup>158</sup>]IFN-αF; and

  [Arg<sup>22</sup>, Ala<sup>76</sup>, Asp<sup>78</sup>, Glu<sup>79</sup>, Tyr<sup>86</sup>, Tyr<sup>90</sup>,

  Leu<sup>96</sup>, Glu<sup>114</sup>, Arg<sup>121</sup>, Thr<sup>156</sup>, Asn<sup>157</sup>, Leu<sup>158</sup>]IFN-αF.
- 39. A manufactured gene capable of directing the synthesis in a selected host microorganism of human leukocyte interferon subtype F.
- 40. A manufactured gene according to claim
  39 wherein the base sequence of the top strand is as
  25 follows:
  5'-TGT-GAT-TTA-CCT-CAA-ACT-CAT-TCT-CTT-GGT-AAC-CGT-CGCGCT-CTG-ATT-CTG-CTG-GCA-CAG-ATG-GGT-CGT-ATT-TTC-CCG-TTTAGC-TGC-CTG-AAA-GAC-CGT-CAC-GAC-TTC-GGC-TTT-CCG-CAA-GAAGAG-TTC-GAT-GGC-AAC-CAA-TTC-CAG-AAA-GCT-CAG-GCA-ATC-TCT30 GTA-CTG-CAC-GAA-ATG-ATC-CAA-CAG-ACC-TTC-AAC-CTG-TTT-TCCACT-AAA-GAC-AGC-TCT-GCT-ACC-TGG-GAA-CAA-AGC-TTG-CTG-GAGAAG-TTC-TCC-ACT-GAA-CTG-AAC-CAG-CAG-CTG-AAC-GAC-ATG-GAAGCA-TGC-GTA-ATC-CAG-GAA-GTT-GGT-GTA-GAA-GAG-ACT-CCG-CTGATG-AAC-GTC-GAC-TCT-ATT-CTG-GCA-GTT-AAA-AAG-TAC-TTC-CAG35 CGT-ATC-ACT-CTG-TAC-CTG-ACC-GAA-AATT-ATG-CGT-TCT-TTC-TCTGCT-TGG-GAA-GTA-GTT-CGC-GCT-GAA-AATT-ATG-CGT-TCT-TTC-TCT-

CTG-AGC-AAA-ATC-TTC-CAG-GAG-CGT-CTG-CGC-CGT-AAA-GAA-3'



41. A process for production of human leukocyte interferon subtype F comprising:

growing, under appropriate nutrient conditions, microorganisms transformed with biologically functional DNA including a manufactured gene according to claim 39, whereby said microorganisms express said gene and produce human leukocyte interferon subtype F.

- 42. The polypeptide product of the process 10 of claim 41.
- 43. A manufactured gene capable of directing the synthesis in a selected host microorganism of a polypeptide which differs from human leukocyte interferon subtype F in terms of the identity and/or location of one or more amino acids.
- 44. A process for production of a polypeptide which differs from human leukocyte interferon subtype
  20 F in terms of the identity and/or location of one or more amino acids comprising:

growing, under appropriate nutrient conditions, microorganisms transformed with biologically functional DNA including a manufactured gene according to claim 43, whereby said microorganisms express said gene and produce said polypeptide.

45. A polypeptide product of the process of claim 44.

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- 46. A polypeptide product according to claim 45 which is [Thr  $^{14}$ , Met  $^{16}$ ] IFN- $\alpha$ F.
- 47. A reagent material comprising a radio-35 labelled manufactured DNA sequence according to claim 6.



- 48. A reagent material according to claim 47 wherein said radiolabel is  $\mathbf{I}^{125}$ .
- 49. A reagent material comprising a tagged 5 antibody to a polypeptide according to claim 12, coated on the surface of a plastic bead.
- 50. A microbially synthesized, biologically active proteinaceous product which differs from a naturally-occurred form thereof in terms of the identity and/or location of one or more amino acids and in terms of one or more biological and pharmacological properties but which substantially retains other such properties of the naturally-occurring form.

51. A polypeptide product according to claim 50 selected from the group consisting of:

[Lys<sup>81</sup>]IFN<sub>\gamma</sub>;

[Ser<sup>1</sup>, Ser<sup>3</sup>, Lys<sup>81</sup>]IFN<sub>\gamma</sub>;

[Lys<sup>1</sup>, Lys<sup>2</sup>, Gln<sup>3</sup>, Lys<sup>81</sup>]IFN<sub>\gamma</sub>;

[des-Cys<sup>1</sup>, des-Tyr<sup>2</sup>, des-Cys<sup>3</sup>, Lys<sup>81</sup>]IFN<sub>\gamma</sub>;

[Phe<sup>39</sup>, Lys<sup>81</sup>]IFN<sub>\gamma</sub>;

[Thr<sup>48</sup>, Lys<sup>81</sup>]IFN<sub>\gamma</sub>;

[Lys<sup>81</sup>, Cys<sup>95</sup>]IFN<sub>\gamma</sub>;

[Cys<sup>95</sup>, Pro<sup>104</sup>]IFN<sub>\gamma</sub>;

[Gln<sup>28</sup>]IFN<sub>\gamma</sub>;

[Gln<sup>28</sup>]IFN<sub>\gamma</sub>;

[Gln<sup>28</sup>]IFN<sub>\gamma</sub>;

[Gln<sup>2</sup>]IFN<sub>\gamma</sub>;

[Gln<sup>2</sup>]IFN<sub>\gamma</sub>;

[Gln<sup>2</sup>]IFN<sub>\gamma</sub>;

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52. A process for enhancing the expression of an exogenous, vector-borne, gene in <a href="E.coli">E.coli</a>, said process comprising:

inserting in said vector, in a position upstream of the exogenous gene protein coding sequence, a DNA sequence comprising base pairs duplicative of



ribosome binding site base pairs associated with  $\underline{\text{E.coli}}$  synthesis of OMP-F, CRO and CAM gene products.

- 53. The process of claim 52 wherein the inserted DNA sequence is a sequence associated with E.coli synthesis of OMP-F gene products.
  - 54. The process of claim 53 wherein the inserted DNA sequence is:

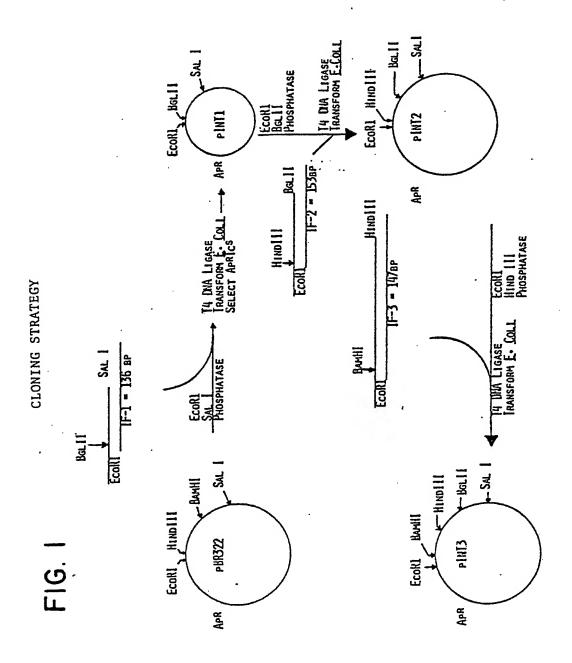
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- 1 10 20 5'-AACCATGAGGGTAATAAATA-3' 3'-TTGGTACTCCCATTATTTAT-5'.
- 55. The process of claim 53 wherein the inserted DNA sequence follows a trp promoter/operator sequence.



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# INTERNATIONAL SEARCH REPORT

International Application No PCT/US83/00605

		International Application No PCI/U	003700003			
I. CLASS	SIFICATION OF SUBJECT MATTER (II several class)	fication symbols apply, indicate all) 3				
TPC:30	g to International Patent Classification (IPC) or to both Nat C12P 19/34; C07H 21/04; C07G 7/00 C12N 15/00; A61K 39/00; A61K 45/0	); C12N 1/00; C12P 21/00	); C12P 21/02; /56			
II. FIELD	S SEARCHED					
	Minimum Docume					
Classificati		Classification Symbols				
435/91,68,70,172,317,253,7; 536/27; 260/112R,112.5R; 424/85, 177; 436/504,513,518,528,531,532,533						
	Documentation Searched other to the Extent that such Documents	than Minimum Documentation are Included in the Fields Searched 5				
r BIOSIS PREVIEWS 1969-83 CA SEARCH 1967-83						
III. DOCL	JMENTS CONSIDERED TO BE RELEVANT 14.					
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Date of the Actual Completion of the International Search 2  16 AUGUST 1983  Date of Mailing of this International Search Report 2  26 AUG 1983						
International Searching Authority 1  ISA/US  Signature of Authorized Officer 80  JAMES MARTINELL						

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VI OBS	ERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 10					
	itional search report has not been established in respect of certain claims under Article 17(2) (a) for	the following reasons:				
	numbers, because they relate to subject matter 12 not required to be searched by this Auti					
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2 Claim	numbers because they relate to parts of the international application that do not comply wi	th the prescribed require-				
ments	to such an extent that no meaningful international search can be carried out 13, specifically:					
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VI.X OBS	ERVATIONS WHERE UNITY OF INVENTION IS LACKING 11					
This Interna	tional Searching Authority found multiple inventions in this international application as follows:					
	required additional search fees were timely paid by the applicant, this international search report cov nternational application.	ers all searchable claims				
	2 As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims of the international application for which fees were paid, specifically claims:					
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	uired additional search fees were timely paid by the applicant. Consequently, this international search trial international search trial mentioned in the claims; it is covered by claim numbers:	ch report is restricted to				
	earchable claims could be searched without effort justifying an additional fee, the International Sea	rching Authority did not				
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